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(71) Applicant: VERTEX PHARMACEUTICALS INCORPO-RATED [US/US]; 40 Allston Street, Cambridge, MA

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(72) Inventors: WILSON, Keith, P.; 6 Longwood Drive, Hopkinton, MA 01748 (US). GRIFFITH, James, P.; 15 Wood Ridge Circle, Weston, MA 02193 (US). KIM, Eunice, E.; 1500 Worcester Road, Framingham, MA 01701 (US). LIV-INGSTON, David, J., 20 Madison Avenue, Newtonville, MA 02160 (US).

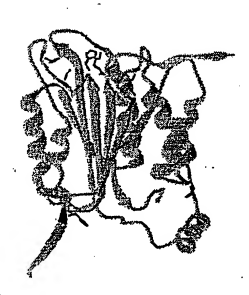
(74) Agents: HALEY, James, F., Jr. et al.; Fish & Neave, 1251 Avenue of the Americas, New York, NY 10020 (US).

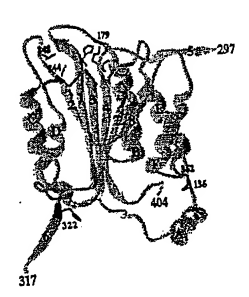
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(54) Tide: CRYSTAL STRUCTURE AND MUTANTS OF INTERLEUKIN-18 CONVERTING ENZYME





(57) Abstract

Interleukin-1\beta converting enzyme ("ICE") processes an inactive precursor to the pro-inflammatory cytokine, interleukin-1\beta. The high-resolution structure of human ICE crystallized in complex with an inhibitor is determined by X-ray diffraction. The active site spans both the 10 and 20 kilodalton subunits. The accessory binding site is composed of residues from the p10 and p20 subunits that are adjacent to the two-fold axis of the crystal. The structure coordinates of the enzyme may be used to design novel classes of ICE inhibitors.

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CRYSTAL STRUCTURE AND MUTANTS OF INTERLEUKIN-16 CONVERTING ENZYME

TECHNICAL FIELD OF INVENTION

The present invention relates to crystals of
interleukin-1ß converting enzyme ("ICE") and more
particularly to the high resolution structure of ICE
obtained by X-ray diffraction. This invention also
relates to mutants of ICE. In addition, this invention
relates to methods of using the structure coordinates
of ICE and mutants thereof to screen and design
compounds that bind to the active site and accessory
binding site of ICE.

BACKGROUND ART

Interleukin-1 ("IL-1") is a major proinflammatory and immunoregulatory protein that
stimulates fibroblast differentiation and
proliferation, the production of prostaglandins,
collagenase and phospholipase by synovial cells and
chondrocytes, basophil and eosinophil degranulation and
neutrophil activation. Oppenheim, J.H. et al,
Immunology Today, 7, pp. 45-56 (1986). As such, it is
involved in the pathogenesis of chronic and acute
inflammatory and autoimmune diseases. IL-1 is
predominantly produced by peripheral blood monocytes
and exists in two distinct agonist forms, IL-1α and IL1β. Mosely, B.S. et al., Proc. Nat. Acad. Sci., 84,

pp. 4572-4576 (1987); Lonnemann G. et al., <u>Eur.J.</u> <u>Immunol.</u>, 19, pp. 1531-1536 (1989).

IL-1ß is synthesized as a biologically inactive precursor, pIL-1ß. pIL-1ß is a 33kDa

5 polypeptide that lacks a conventional leader sequence and is not processed by a signal peptidase. March, C.J., Nature, 315, pp. 641-647 (1985). Instead, pIL-1ß is cleaved by interleukin-1ß converting enzyme ("ICE") between Asp 116 and Ala 117 to produce the biologically active C-terminal fragment of 17kDa molecular weight found in serum and synovial fluid. Sleath, P.R. et al., J. Biol. Chem., 265, pp. 14526-14528 (1992); Howard, A.D. et al., J. Immunol., 147, pp. 2964-2969 (1991). Processing by ICE is also necessary for the transport of mature IL-1ß through the cell membrane.

ICE is a cysteine protease localized primarily in monocytes. It converts precursor IL-16 to the mature form. Black, R.A. et al., <u>FEBS Lett.</u>, 247, pp. 386-390 (1989); Kostura, M.J. et al., <u>Proc. Natl.</u>

- Acad. Sci. USA, 86, pp. 5227-5231 (1989). ICE, or its homologues, also appears to be involved in the regulation of cell death or apoptosis. Yuan, J. et al., Cell, 75, pp. 641-652 (1993); Miura, M. et al., Cell, 75, pp. 653-660 (1993); Nett-Fiordalisi, M.A.
- et al., J. Cell Biochem., 17B, p. 117 (1993). In particular, ICE or ICE homologues are thought to be associated with the regulation of apoptosis in neurogenerative diseases, such as Alzheimer's and Parkinson's disease. Marx, J. and M. Baringa, Science,
- 30 259, pp. 760-762 (1993); Gagliardini, V. et al., Science, 263, pp. 826-828 (1994).

ICE has been previously described as a heterodimer composed of two subunits, p20 and p10 (20kDa and 10kDa molecular weight, respectively).

35 These subunits are derived from a 45kDa proenzyme (p45)

- 3 -

by way of a p30 form, through an activation mechanism that is autocatalytic. Thornberry, N.A. et al.,

Nature, 356, pp. 768-774 (1992). The ICE proenzyme has been divided into several functional domains: a

prodomain (p14), a p22/20 subunit, a polypeptide linker and a p10 subunit. Thornberry et al., supra; Casano et al., Genomics, 20, pp. 474-481 (1994).

Full length p45 has been characterized by its cDNA and amino acid sequences. PCT patent applications WO 91/15577 and WO 94/00154. The p20 and p10 cDNA and amino acid sequences are also known. Thornberry et al., supra. Murine and rat ICE have also been sequenced and cloned. They have high amino acid and nucleic acid sequence homology to human ICE. Miller, D.K. et al., Ann. N.Y. Acad. Sci., 696, pp. 133-148

(1993); Molineaux, S.M. et al., <u>Proc. Nat. Acad. Sci.</u>, 90, pp. 1809-1813 (1993). Knowledge of the primary structure of ICE, however, does not allow prediction of its tertiary structure. Nor does it afford an

20 understanding of the structural, conformational and chemical interactions of ICE and its substrate pIL-18 or other substrates or inhibitors.

ICE inhibitors represent a class of compounds useful for the control of inflammation or apoptosis or both. Peptide and peptidyl inhibitors of ICE have been described. PCT patent applications WO 91/15577; WO 93/05071; WO 93/09135; WO 93/14777 and WO 93/16710; and European patent application 0 547 699. However, due to their peptidic nature, such inhibitors are typically characterized by undesirable pharmacologic properties, such as poor oral absorption, poor stability and rapid metabolism. Plattner, J.J. and D.W. Norbeck, in Drug Discovery Technologies, C.R. Clark and W.H. Moos, Eds. (Ellis Horwood, Chichester, England, 1990), pp. 92-

- 4 -

126. This has hampered their development into effective drugs.

SUMMARY OF THE INVENTION

The present invention solves the above problems.

It is an object of this invention to solve the three-dimensional structure of interleukin-1ß converting enzyme ("ICE") and to determine its structure coordinates.

It is an object of this invention to use the structure coordinates of an ICE crystal to reveal the atomic details of the active site and one or more accessory binding sites of the enzyme.

It is also an object of this invention to use the structure coordinates of an ICE crystal to solve the structure of a different ICE crystal, or a crystal of a mutant, homologue or co-complex, of ICE.

It is a further object of this invention to provide interleukin-16 converting enzyme mutants

characterized by one or more different properties as compared with wild-type ICE. These properties include altered surface charge, increased stability to subunit dissociation, altered substrate specificity or higher specific activity. ICE mutants are useful to identify those amino acids that are most important for the enzymatic activity of ICE. This information, in turn, allows the design of improved inhibitors of ICE as compared with peptidic ICE inhibitors.

It is also an object of this invention to use the structure coordinates and atomic details of ICE, or its mutants or homologues or co-complexes, to design, evaluate computationally, synthesize and use inhibitors of ICE that avoid the undesirable physical and pharmacologic properties of peptidic ICE inhibitors.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 represents a ribbon drawing of the p20/p10 interleukin-18 converting enzyme heterodimer.

The active site is at the top of the figure, roughly at the center of the cluster of displayed side chains.

Figure 2 represents a space-filling model of the (p20)₂/(p10)₂ tetramer of interleukin-1ß converting enzyme. Two p20 subunits (dark shade) are in contact with two p10 subunits (light shade). Black shading on top left and bottom right represents a tetrapeptide aldehyde inhibitor bound in each of the two active sites of the tetramer. The crystallographic two-fold axis is approximately perpendicular to the plane of drawing, and runs through the small hole at the center of the interface between the two p10 subunits. The N-and C-terminal ends of each subunit are labeled.

Figure 3 is a graphic depiction of the activity of various interleukin-1ß converting enzyme mutants in processing pIL-1ß intracellularly, relative to wild-type interleukin-1ß converting enzyme activity. The particular mutants tested are designated on the x-axis using nomenclature listing the specific amino acid and its residue number. For example, "C285S" indicates replacement of amino acid Cys-285 with serine. Activity levels were measured at 16 hours (hatched bar) and 24 hours (solid bars).

BRIEF DESCRIPTION OF THE TABLES

Table A lists the amino acids of ICE that constitute the tetramer interface contacts between the ICE subunits and that constitute the accessory binding site moiety.

Table B lists the atomic structure coordinates for ICE as derived by X-ray diffraction

- 6 -

from a crystal of ICE complexed to a tetrapeptide inhibitor.

ABBREVIATIONS AND DEFINITIONS

ABBREVIATIONS

5 Amino Acids

A = Ala = Alanine

V = Val = Valine

L = Leu = Leucine

I = Ile = Isoleucine

10 P = Pro = Proline

F = Phe = Phenylalanine

W = Trp = Tryptophan

M = Met = Methionine

G = Gly = Glycine

15 S = Ser = Serine

T = Thr = Threonine

C = Cys = Cysteine

Y = Tyr = Tyrosine

N = Asn = Asparagine

20 Q = Gln = Glutamine

D = Asp = Aspartic Acid

E = Glu = Glutamic Acid

K = Lys = Lysine

R = Arg = Arginine

25 H = His = Histidine

. DEFINITIONS

The following terms are also used herein:

The term "naturally occurring amino acids"

means the L-isomers of the naturally occurring amino

acids. The naturally occurring amino acids are
glycine, alanine, valine, leucine, isoleucine, serine,

methionine, threonine, phenylalanine, tyrosine,
tryptophan, cysteine, proline, histidine, aspartic

- 7 -

acid, asparagine, glutamic acid, glutamine, γ -carboxyglutamic acid, arginine, ornithine and lysine. Unless specifically indicated, all amino acids referred to in this application are in the <u>L</u>-form.

The term "unnatural amino acids" means amino acids that are not naturally found in proteins.

Examples of unnatural amino acids used herein, include racemic mixtures of selenocysteine and selenomethionine. In addition, unnatural amino acids include the D or L forms of nor-leucine, paranitrophenylalanine, homophenylalanine, parafluorophenylalanine, 3-amino-2-benzylpropionic acid, homoarginine, and D-phenylalanine.

The term "positively charged amino acid"

includes any naturally occurring or unnatural amino acid having a positively charged side chain under normal physiological conditions. Examples of positively charged naturally occurring amino acids are arginine, lysine and histidine.

The term "negatively charged amino acid" includes any naturally occurring or unnatural amino acid having a negatively charged side chain under normal physiological conditions. Examples of negatively charged naturally occurring amino acids are aspartic acid and glutamic acid.

The term "hydrophobic amino acid" means any amino acid having an uncharged, nonpolar side chain that is relatively insoluble in water. Examples of naturally occurring hydrophobic amino acids are alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine.

The term "hydrophilic amino acid" means any amino acid having an uncharged, polar side chain that is relatively soluble in water. Examples of naturally occurring hydrophilic amino acids are serine, infra,

threonine, tyrosine, asparagine, glutamine, and cysteine.

The term "mutant" refers to an ICE
polypeptide, i.e.. a polypeptide displaying the

5 biological activity of wild-type, human ICE,
characterized by the replacement of at least one amino
acid from the wild-type, human ICE sequence according
to Thornberry, N.A. et al., Nature, 356, pp. 768-774
(1992). Such a mutant may be prepared, for example, by
expression of ICE cDNA previously altered in its coding
sequence by oligonucleotide-directed mutagenesis.

specific incorporation of unnatural amino acids into ICE proteins using the general biosynthetic method of Noren, C.J., et al., Science, 244, pp. 182-188 (1989). In this method, the codon encoding the amino acid of interest in wild-type ICE is replaced by a "blank" nonsense codon, TAG, using oligonucleotide-directed mutagenesis (described in detail, infra). A suppressor tRNA directed against this codon is then chemically aminoacylated in vitro with the desired unnatural amino acid. The aminoacylated tRNA is then added to an in vitro translation system to yield a mutant ICE enzyme with the site-specific incorporated unnatural amino acid.

Selenocysteine or selenomethionine may be incorporated into wild-type or mutant ICE by expression of ICE-encoding cDNAs in auxotrophic <u>E. coli</u> strains. Hendrickson, W.A. et al., <u>EMBO J.</u>, 9(5), pp. 1665-1672 (1990). In this method, the wild-type or mutagenized ICE cDNA may be expressed in a host organism on a growth medium depleted of either natural cysteine or methionine (or both) but enriched in selenocysteine or selenomethionine (or both).

25

The term "altered surface charge" means a change in one or more of the charge units of a mutant polypeptide, at physiological pH, as compared to wildtype ICE. This is preferably achieved by mutation of 5 at least one amino acid of wild-type ICE to an amino acid comprising a side chain with a different charge at physiological pH than the original wild-type side chain.

The change in surface charge is determined by 10 measuring the isoelectric point (pI) of the polypeptide molecule containing the substituted amino acid and comparing it to the isoelectric point of the wild-type ICE molecule.

The term "high specific activity" refers to a specific activity of ICE where the second-order rate constant (k_{cat}/K_m) for hydrolysis of the substrate Ac-Tyr-Val-Ala-Asp-aminomethylcoumarin exceeds 7 x 104 M⁻¹s⁻¹ at 25°C, using the assay described by Pennington, M.W. and N.A. Thornberry, Peptide Res., 7(2), pp. 72-20 76 (1994). Alternatively, the specific activity of ICE may be determined by monitoring hydrolysis of the substrate Ac-Tyr-Val-Ala-Asp-p-nitroaniline. Reiter, L.A., Intr. J. Peptide Protein Res., 43, pp. 8796 (1994).

The term "altered substrate specificity" refers to a change in the ability of a mutant ICE to cleave a substrate as compared to wild-type ICE. Substrate specificity may be measured by hydrolysis of fluorogenic peptide substrates or of unmodified peptide 30 substrates by ICE, as described in Thornberry et al., supra. ICE mutants with altered substrate specificity demonstrate a second order rate constant (k_{cat}/K_m) for a substrate X_1 -Tyr-Val-Ala- X_2 - X_3 that exceeds the k_{cat}/K_m for the analogous peptide substrate, X1-Tyr-Val-Ala-35 Asp-X3. For both substrates, X1 is an amino protecting

PCT/US95/07619 WO 95/35367

- 10 -

group, such as acetyl; X2 is a natural or unnatural amino acid residue other than L-aspartate; and X_3 is a carboxyl protecting group, such as aminomethylcoumarin or p-nitroaniline.

The "kinetic form" of ICE refers to the condition of the enzyme in its free or unbound form or bound to a chemical entity at either its active site or accessory binding site.

A "competitive" inhibitor is one that 10 inhibits ICE activity by binding to the same kinetic form, of ICE, as its substrate binds -- thus directly competing with the substrate for the active site of ICE. Competitive inhibition can be reversed completely by increasing the substrate concentration.

15

An "uncompetitive" inhibitor is one that inhibits ICE by binding to a different kinetic form of the enzyme than does the substrate. Such inhibitors bind to ICE already bound with the substrate and not to the free enzyme. Uncompetitive inhibition cannot be 20 reversed completely by increasing the substrate concentration.

A "non-competitive" inhibitor is one that can bind to either the free or substrate bound form of ICE.

Those of skill in the art may identify 25 inhibitors as competitive, uncompetitive or noncompetitive, by computer fitting enzyme kinetic data using standard equations according to Segel, I.H., Enzyme Kinetics, J. Wiley & Sons, (1975). It should also be understood that uncompetitive or non-30 competitive inhibitors according to this invention may bind to the accessory binding site.

The term "homologue" means a protein having at least 30% amino acid sequence identity with ICE or any functional domain of ICE as defined by Thornberry 35 et al., supra and Casano et al., supra.

- 11 -

The term "subunit dissociation" refers to the fact that at very high dilutions of wild-type ICE, or at concentrations of the enzyme below 10 nM, enzymatic activity shows a time dependent loss assayed in the 5 presence of a tetrapeptide substrate. Reconcentration of the dilute, inactive mixture results in complete recovery of ICE activity. Wild-type ICE demonstrates a Kd for subunit dissociation between 1 and 10 nM. Enzymatic activity is determined by measuring the activity of ICE according to the assay of Pennington and Thornberry, supra, at varying concentrations of the enzyme. The concentration of the enzyme is determined by active site titration.

The term "co-complex" means ICE or a mutant or homologue of ICE in covalent or non-covalent association with a chemical entity or compound.

The term "associating with" refers to a condition of proximity between a chemical entity or compound, or portions thereof, and an ICE molecule or portions thereof. The association may be non-covalent -- wherein the juxtaposition is energetically favored by hydrogen bonding or van der Waals or electrostatic interactions -- or it may be covalent.

The term "G-sheet" refers to the conformation

of a polypeptide chain stretched into an extended zigzig conformation. Portions of polypeptide chains that
run "parallel" all run in the same direction.

Polypeptide chains that are "antiparallel" run in the
opposite direction from the parallel chains.

The term "active site" or "active site
moiety" refers to any or all of the following sites in
ICE: the substrate binding site; the site where the
tetrapeptide inhibitor binds and the site where the
cleavage of a substrate occurs. The active site is
characterized by at least amino acid residues 173, 176,

PCT/US95/07619 WO 95/35367

- 12 -

177, 178, 179, 180, 236, 237, 238, 239, 244, 248, 283, 284, 285, 290, 338, 339, 340, 341, 342, 343, 344, 345, 348, 352, 381 and 383, using the sequence and numbering according to Thornberry et al., supra (SEQ ID NO:1).

The term "accessory binding site" or "accessory binding site moiety" refers to a binding site of ICE comprising amino acid residues adjacent to the two-fold axis of ICE but external to the active site, according to Table A. An accessory binding site 10 may be a locus of ICE inhibition, although it is not the site of substrate cleavage.

The accessory binding site is characterized by at least amino acid residues 150, 151, 240, 259, 267, 268, 274, 291, 292, 293, 294, 295, 296, 297, 317, 15 318, 319, 320, 321, 322, 323, 324, 325, 327, 334, 335, 367, 371, 374, 375, 377, 378, 380, 382, 384, 386, 388, 389, 390, 391, 392, 393, 394, 395 and 396, using the sequence and numbering according to Thornberry et al., supra (SEQ ID NO:1).

20

The term "P binding pocket" refers to a binding subsite, or portion of the binding site on the ICE molecule. The amino acid residues of an ICE substrate are given designations according to their position relative to the scissile bond, i.e. the bond 25 that is broken by the protease. Residues are designated P1, P2, etc., for those extending toward the N-terminus from the scissile bond of the substrate. The residues are designated P1', P2', etc., for those extending toward the C-terminus from the scissile bond 30 of the substrate.

The portions of an ICE inhibitor that correspond to the P or P' residues of the substrate are also labeled P1, P1', etc., by analogy with the substrate. The binding subsites of the ICE molecule 35 that receive the residues labeled P1, P1', etc., are

- 13 -

designated "the S1 site", "the P1' binding pocket", etc. Schechter, I. and A. Berger, "On the Size of the Active Site in Proteases", <u>Biochem. Biophys. Res.</u>
Commun., 27, pp. 157-162 (1967).

The "P1 binding pocket" of the ICE active site is defined as the space surrounded by amino acid residues Arg-179, His-237, Gln-283 and Arg-341.

The "P2 binding pocket" of the ICE active site is defined as the space surrounded by amino acid residues Pro-290, Val-338 and Trp-340.

The "P3 binding pocket" of the ICE active site is defined as the space surrounded by amino acid residues Pro-177, Arg-178, Thr-180, Arg-341 and Pro-343.

The "P4 binding pocket" of the ICE active site is defined as the space surrounded by amino acid residues Trp-340, His-342, Met-345, Val-348, Arg-352, Asp-381 and Arg-383.

The "P' binding pocket" of the ICE active 20 site is defined as the space surrounded by amino acid residues Phe-173, Ile-176, His-237, Gly-238, Ile-239, Cys-244 and His-248.

The term "pl0 subunits interacting across the two-fold axis" means having at least 50% of the interface contacts according to Table A.

The term "structure coordinates" refers to mathematical coordinates derived from mathematical equations related to the patterns obtained on diffraction of a monochromatic beam of X-rays by the atoms (scattering centers) of an ICE molecule in crystal form. The diffraction data are used to calculate an electron density map of the repeating unit of the crystal. The electron density maps are used to establish the positions of the individual atoms within the unit cell of the crystal.

The term "heavy atom derivatization" refers to the method of producing a chemically modified form of a crystal of ICE. In practice, a crystal is soaked in a solution containing heavy metal atom salts, or organometallic compounds, e.g., lead chloride, gold thiomalate, thimerosal or uranyl acetate, which can diffuse through the crystal and bind to the surface of the protein. The location(s) of the bound heavy metal atom(s) can be determined by X-ray diffraction analysis of the soaked crystal. This information, in turn, is used to generate the phase information used to construct three-dimensional structure of the enzyme. Blundel, T.L. and N.L. Johnson, Protein Crystallography, Academic Press (1976).

15 Those of skill in the art understand that a set of structure coordinates determined by X-ray crystallography is not without standard error. For the purpose of this invention, any set of structure coordinates for ICE or ICE homologues or ICE mutants that have a root mean square deviation of protein backbone atoms (N, Ca, C and O) of less than 0.75Å when superimposed -- using backbone atoms -- on the structure coordinates listed in Table B shall be considered identical.

25 The term "unit cell" refers to a basic parallelipiped shaped block. The entire volume of a crystal may be constructed by regular assembly of such blocks. Each unit cell comprises a complete representation of the unit of pattern, the repetition of which builds up the crystal.

The term "space group" refers to the arrangement of symmetry elements of a crystal.

The term "molecular replacement" refers to a method that involves generating a preliminary model of an ICE crystal whose structure coordinates are unknown,

- 15 -

by orienting and positioning a molecule whose structure coordinates are known (e.g., ICE coordinates from Table B) within the unit cell of the unknown crystal so as best to account for the observed diffraction pattern of 5 the unknown crystal. Phases can then be calculated from this model and combined with the observed amplitudes to give an approximate Fourier synthesis of the structure whose coordinates are unknown. This, in turn, can be subject to any of the several forms of 10 refinement to provide a final, accurate structure of the unknown crystal. Lattman, E., *Use of the Rotation and Translation Functions", in Methods in Enzymology, 115, pp. 55-77 (1985); M.G. Rossmann, ed., "The Molecular Replacement Method", Int. Sci. Rev. Ser., 15 No. 13, Gordon & Breach, New York, (1972). Using the structure coordinates of ICE provided by this invention, molecular replacement may be used to determine the structure coordinates of a crystalline mutant or homologue of ICE or of a different crystal 20 form of ICE.

DETAILED DESCRIPTION OF THE INVENTION

In order that the invention described herein may be more fully understood, the following detailed description is set forth.

The present invention relates to crystalline 25 interleukin-16 converting enzyme ("ICE"), the structure of ICE as determined by X-ray crystallography, the use of that structure to solve the structure of ICE homologues and of other crystal forms of ICE, mutants 30 and co-complexes of ICE, and the use of the ICE structure and that of its homologues, mutants and cocomplexes to design inhibitors of ICE.

PCT/US95/07619 WO 95/35367

- 16 -

The Structure of ICE Α.

The present invention provides, for the first time, crystals of human ICE grown in the presence of a tetrapeptide inhibitor from solutions of polyethylene 5 glycol, as well as the structure of ICE as determined therefrom. The crystals have tetragonal space group symmetry P43212. The unit cell of said crystals has a rectangular shape of dimensions a=b=65 ± 5Å, and c=162 ± 5Å. The structure coordinates of ICE, as 10 determined by X-ray crystallography of crystalline ICE, are listed in Table B.

Crystal packing reveals that ICE is a $(p20)_2/(p10)_2$ tetramer. In the tetramer, two p20 subunits contact two adjacent p10 subunits which 15 interact across the crystallographic two-fold axis (Figure 2). This axis corresponds to an oligomer interface in solution. Most of the dimer-dimer interface consists of p20 residues 291-297 and of p10 residues 318-322 and 386-396.

Figure 1 represents a ribbon drawing of the 20 p20/p10 ICE heterodimer. As depicted in the figure, the p20 and p10 subunits are intimately associated and the active site is at the top of the figure, roughly at the center of the cluster of displayed side chains.

25

The enzyme core is a six-stranded B-sheet with 5 parallel strands (numbered 1, 2, 3, 4 and 7) and one anti-parallel strand (numbered 8). Six α -helices (lettered A, B, C, D, E and F) lie roughly parallel to the B-strands. The last seven residues of p20 and the 30 first seven of pl0 protrude from this compact structure and form two anti-parallel B-strands [5 (residues 291-297)] and 6 (residues 317-323)]. A few key residues are labelled according to their position in the p45 amino acid sequence of ICE (Thornberry et al., supra).

- 17 -

Our understanding of the structure of ICE has enabled, for the first time, identification of the active site and accessory binding site of the enzyme. The p10 subunit from one ICE molecule contacts the p20 5 subunit from a different molecule and together they create an active site. The active site spans both the p20 and p10 subunits and comprises amino acid residues from both subunits. The active site moiety is characterized by at least amino acid residues 173, 176, 10 177, 178, 179, 180, 236, 237, 238, 239, 244, 248, 283, 284, 285, 290, 338, 339, 340, 341, 342, 343, 344, 345, 348, 352, 381 and 383 using the sequence numbering according to Thornberry et al., supra (SEQ ID NO:1). An accessory binding site is formed by amino

15 acid residues on the p10 subunits that interact across the two-fold axis. The accessory binding site moiety is characterized by at least amino acid residues 150, 151, 240, 259, 267, 268, 274, 291, 292, 293, 294, 295, 296, 297, 317, 318, 319, 320, 321, 322, 323, 324, 325, 20 327, 334, 335, 367, 371, 374, 375, 377, 378, 378, 380, 382, 384, 386, 388, 389, 390, 391, 392, 393, 394, 395 and 396 using the sequence numbering according to Thornberry et al., supra (SEQ ID NO:1).

Uses of the Structure Coordinates of ICE В.

For the first time, the present invention 25 permits the use of molecular design techniques to design, select and synthesize chemical entities and compounds, including inhibitory compounds, capable of binding to the active site or accessory binding site of 30 ICE, in whole or in part.

On approach enabled by this invention, is to use the structure coordinates of ICE to design compounds that bind to the enzyme and alter the physical properties of the compounds in different ways,

- 18 -

e.g., solubility. For example, this invention enables the design of compounds that act as competitive inhibitors of the ICE enzyme by binding to, all or a portion of, the active site of ICE. This invention
5 also enables the design of compounds that act as uncompetitive inhibitors of the ICE enzyme. These inhibitors may bind to, all or a portion of, the accessory binding site of an ICE already bound to its substrate and may be more potent and less non-specific
10 than known competitive inhibitors that compete only for the ICE active site. Similarly, non-competitive inhibitors that bind to and inhibit ICE whether or not it is bound to another chemical entity may be designed using the structure coordinates of ICE of this
15 invention.

A second design approach is to probe an ICE crystal with molecules composed of a variety of different chemical entities to determine optimal sites for interaction between candidate ICE inhibitors and the enzyme. For example, high resolution X-ray diffraction data collected from crystals saturated with solvent allows the determination of where each type of solvent molecule sticks. Small molecules that bind tightly to those sites can then be designed and synthesized and tested for their ICE inhibitor activity. Travis, J., Science, 262, p. 1374 (1993).

This invention also enables the development of compounds that can isomerize to short-lived reaction intermediates in the chemical reaction of a substrate or other compound that binds to ICE, with ICE. Thus, the time-dependent analysis of structural changes in ICE during its interaction with other molecules is enabled. The reaction intermediates of ICE can also be deduced from the reaction product in co-complex with ICE. Such information is useful to design improved

- 19 -

analogues of known ICE inhibitors or to design novel classes of inhibitors based on the reaction intermediates of the ICE enzyme and ICE-inhibitor co-complex. This provides a novel route for designing ICE inhibitors with both high specificity and stability.

Another approach made possible and enabled by this invention, is to screen computationally small molecule data bases for chemical entities or compounds that can bind in whole, or in part, to the ICE enzyme.

10 In this screening, the quality of fit of such entities or compounds to the binding site may be judged either by shape complementarity or by estimated interaction energy. Meng, E.C. et al., <u>J. Comp. Chem.</u>, 13, pp. 505-524 (1992).

Because ICE may crystallize in more than one crystal form, the structure coordinates of ICE, or portions thereof, as provided by this invention are particularly useful to solve the structure of those other crystal forms of ICE. They may also be used to solve the structure of ICE mutants, ICE co-complexes, or of the crystalline form of any other protein with significant amino acid sequence homology to any functional domain of ICE.

One method that may be employed for this
purpose is molecular replacement. In this method, the
unknown crystal structure, whether it is another
crystal form of ICE, an ICE mutant, or an ICE cocomplex, or the crystal of some other protein with
significant amino acid sequence homology to any
functional domain of ICE, may be determined using the
ICE structure coordinates of this invention as provided
in Table B. This method will provide an accurate
structural form for the unknown crystal more quickly
and efficiently than attempting to determine such
information ab initio.

WO 95/35367

In addition, in accordance with this invention, ICE mutants may be crystallized in cocomplex with known ICE inhibitors. The crystal structures of a series of such complexes may then be solved by molecular replacement and compared with that of wild-type ICE. Potential sites for modification within the various binding sites of the enzyme may thus be identified. This information provides an additional tool for determining the most efficient binding interactions, for example, increased hydrophobic interactions, between ICE and a chemical entity or compound.

All of the complexes referred to above may be studied using well-known X-ray diffraction techniques

15 and may be refined versus 2-3Å resolution X-ray data to an R value of about 0.20 or less using computer software, such as X-PLOR (Yale University, ©1992, distributed by Molecular Simulations, Inc.). See, e.g., Blundel & Johnson, supra; Methods in Enzymology, vol. 114 & 115, H.W. Wyckoff et al., eds., Academic Press (1985). This information may thus be used to optimize known classes of ICE inhibitors, and more importantly, to design and synthesize novel classes of ICE inhibitors.

25 The structure coordinates of ICE mutants provided in this invention also facilitate the identification of related proteins or enzymes analogous to ICE in function, structure or both, thereby further leading to novel therapeutic modes for treating or preventing IL-1 mediated diseases.

The design of compounds that bind to or inhibit ICE according to this invention generally involves consideration of two factors. First, the compound must be capable of physically and structurally associating with ICE. Non-covalent molecular

interactions important in the association of ICE with its substrate include hydrogen bonding, van der Waals and hydrophobic interactions.

Second, the compound must be able to assume a conformation that allows it to associate with ICE. Although certain portions of the compound will not directly participate in this association with ICE, those portions may still influence the overall conformation of the molecule. This, in turn, may have a significant impact on potency. Such conformational requirements include the overall three-dimensional structure and orientation of the chemical entity or compound in relation to all or a portion of the binding site, e.g., active site or accessory binding site of ICE, or the spacing between functional groups of a compound comprising several chemical entities that directly interact with ICE.

The potential inhibitory or binding effect of a chemical compound on ICE may be analyzed prior to its actual synthesis and testing by the use of computer modelling techniques. If the theoretical structure of the given compound suggests insufficient interaction and association between it and ICE, synthesis and testing of the compound is obviated. However, if computer modelling indicates a strong interaction, the molecule may then be synthesized and tested for its ability to bind to ICE and inhibit using the fluorescent substrate assay of Thornberry et al., supra. In this manner, synthesis of inoperative compounds may be avoided.

An inhibitory or other binding compound of ICE may be computationally evaluated and designed by means of a series of steps in which chemical entities or fragments are screened and selected for their

- 22 -

ability to associate with the individual binding pockets or other areas of ICE.

One skilled in the art may use one of several methods to screen chemical entities or fragments for their ability to associate with ICE and more particularly with the individual binding pockets of the ICE active site or accessory binding site. This process may begin by visual inspection of, for example, the active site on the computer screen based on the ICE coordinates in Table B. Selected fragments or chemical entities may then be positioned in a variety of orientations, or docked, within an individual binding pocket of ICE as defined supra. Docking may be accomplished using software such as Quanta and Sybyl, followed by energy minimization and molecular dynamics with standard molecular mechanics forcefields, such as CHARMM and AMBER.

Specialized computer programs may also assist in the process of selecting fragments or chemical

20 entities. These include:

25

- 1. GRID (Goodford, P.J., "A Computational Procedure for Determining Energetically Favorable Binding Sites on Biologically Important Macromolecules", J. Med. Chem., 28, pp. 849-857 (1985)). GRID is available from Oxford University, Oxford, UK.
- 2. MCSS (Miranker, A. and M. Karplus, "Functionality Maps of Binding Sites: A Multiple Copy Simultaneous Search Method." Proteins:

 Structure, Function and Genetics, 11, pp. 29-34

 (1991)). MCSS is available from Molecular Simulations, Burlington, MA.
- 3. AUTODOCK (Goodsell, D.S. and A.J. Olsen,
 "Automated Docking of Substrates to Proteins by
 Simulated Annealing", <u>Proteins: Structure</u>,
 Function, and Genetics, 8, pp. 195-202 (1990)).
 AUTODOCK is available from Scripps Research
 Institute, La Jolla, CA.
- DOCK (Kuntz, I.D. et al., "A Geometric Approach to Macromolecule-Ligand Interactions", <u>J. Mol.</u>
 Biol., 161, pp. 269-288 (1982)). DOCK is

- 23 -

available from University of California, San Francisco, CA.

Once suitable chemical entities or fragments have been selected, they can be assembled into a single compound or inhibitor. Assembly may be proceed by visual inspection of the relationship of the fragments to each other on the three-dimensional image displayed on a computer screen in relation to the structure coordinates of ICE. This would be followed by manual model building using software such as Quanta or Sybyl.

Useful programs to aid one of skill in the art in connecting the individual chemical entities or fragments include:

- 1. CAVEAT (Bartlett, P.A. et al, "CAVEAT: A Program to Facilitate the Structure-Derived Design of Biologically Active Molecules". In Molecular Recognition in Chemical and Biological Problems", Special Pub., Royal Chem. Soc., 78, pp. 182-196 (1989)). CAVEAT is available from the University of California, Berkeley, CA.
 - 2. 3D Database systems such as MACCS-3D (MDL Information Systems, San Leandro, CA). This area is reviewed in Martin, Y.C., "3D Database Searching in Drug Design", <u>J. Med. Chem.</u>, 35, pp. 2145-2154 (1992)).
 - 3. HOOK (available from Molecular Simulations, Burlington, MA).

25

Instead of proceeding to build an ICE inhibitor in a step-wise fashion one fragment or chemical entity at a time as described above, inhibitory or other ICE binding compounds may be designed as a whole or "de novo" using either an empty active site or optionally including some portion(s) of a known inhibitor(s). These methods include:

35 1. LUDI (Bohm, H.-J., "The Computer Program LUDI: A New Method for the De Novo Design of Enzyme Inhibitors", <u>J. Comp. Aid. Molec. Design</u>, 6, pp. 61-78 (1992)). LUDI is available from Biosym Technologies, San Diego, CA.

- LEGEND (Nishibata, Y. and A. Itaï, <u>Tetrahedron</u>, 47, p. 8985 (1991)). LEGEND is available from Molecular Simulations, Burlington, MA.
- LeapFrog (available from Tripos Associates,
 St. Louis, MO).

Other molecular modelling techniques may also be employed in accordance with this invention. See, e.g., Cohen, N.C. et al., "Molecular Modeling Software and Methods for Medicinal Chemistry, J. Med. Chem., 33,

10 pp. 883-894 (1990). See also, Navia, M.A. and M.A.
 Murcko, "The Use of Structural Information in Drug
 Design", Current Opinions in Structural Biology, 2,
 pp. 202-210 (1992).

Once a compound has been designed or selected by the above methods, the efficiency with which that compound may bind to ICE may be tested and optimized by computational evaluation. For example, a compound that has been designed or selected to function as an ICE-inhibitor must also preferably traverse a volume not overlapping that occupied by the active site when it is bound to the native substrate. An effective ICE inhibitor must preferably demonstrate a relatively small difference in energy between its bound and free states (i.e., a small deformation energy of binding).

25 Thus, the most efficient ICE inhibitors should preferably be designed with a deformation energy of binding of not greater than about 10 kcal/mole, preferably, not greater than 7 kcal/mole. ICE inhibitors may interact with the enzyme in more than one conformation that is similar in overall binding energy. In those cases, the deformation energy of binding is taken to be the difference between the energy of the free compound and the average energy of the conformations observed when the inhibitor binds to

35 the enzyme.

- 25 -

A compound designed or selected as binding to ICE may be further computationally optimized so that in its bound state it would preferably lack repulsive electrostatic interaction with the target enzyme. Such non-complementary (e.g., electrostatic) interactions include repulsive charge-charge, dipole-dipole and charge-dipole interactions. Specifically, the sum of all electrostatic interactions between the inhibitor and the enzyme when the inhibitor is bound to ICE, preferably make a neutral or favorable contribution to the enthalpy of binding.

Specific computer software is available in the art to evaluate compound deformation energy and electrostatic interaction. Examples of programs

15 designed for such uses include: Gaussian 92, revision C [M.J. Frisch, Gaussian, Inc., Pittsburgh, PA ©1992];

AMBER, version 4.0 [P.A. Kollman, University of California at San Francisco, ©1994]; QUANTA/CHARMM [Molecular Simulations, Inc., Burlington, MA ©1994];

20 and Insight II/Discover (Biosysm Technologies Inc., San Diego, CA ©1994). These programs may be implemented, for instance, using a Silicon Graphics workstation, IRIS 4D/35 or IBM RISC/6000 workstation model 550. Other hardware systems and software packages will be known to those skilled in the art.

Once an ICE-binding compound has been optimally selected or designed, as described above, substitutions may then be made in some of its atoms or side groups in order to improve or modify its binding properties. Generally, initial substitutions are conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the original group. It should, of course, be understood that components known in the art to alter conformation should be avoided. Such substituted

chemical compounds may then be analyzed for efficiency of fit to ICE by the same computer methods described in detail, above.

C. Mutants Of ICE

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The present invention also enables mutants of ICE and the solving of their crystal structure. More particularly, by virtue of the present invention, the location of the active site, accessory binding site and interface of ICE based on its crystal structure permits the identification of desirable sites for mutation.

For example, mutation may be directed to a particular site or combination of sites of wild-type ICE, i.e., the accessory binding site or only the active site, or a location on the interface site may be chosen for mutagenesis. Similarly, only a location on, at or near the enzyme surface may be replaced, resulting in an altered surface charge of one or more charge units, as compared to the wild-type enzyme. Alternatively, an amino acid residue in ICE may be chosen for replacement based on its hydrophilic or hydrophobic characteristics.

Such mutants may be characterized by any one of several different properties as compared with wild-type ICE. For example, such mutants may have altered surface charge of one or more charge units, or have an increased stability to subunit dissociation. Or such mutants may have an altered substrate specificity in comparison with, or a higher specific activity than, wild-type ICE.

The mutants of ICE prepared by this invention may be prepared in a number of ways. For example, the wild-type sequence of ICE may be mutated in those sites identified using this invention as desirable for mutation, by means of oligonucleotide-directed

- 27 -

deletion. Alternatively, mutants of ICE may be generated by the site specific replacement of a particular amino acid with an unnaturally occurring amino acid. In addition, ICE mutants may be generated through replacement of an amino acid residue, or a particular cysteine or methionine residue, with selenocysteine or selenomethionine. This may be achieved by growing a host organism capable of expressing either the wild-type or mutant polypeptide on a growth medium depleted of either natural cysteine or methionine (or both) but enriched in selenocysteine or selenomethionine (or both).

Mutations may be introduced into a DNA

15 sequence coding for ICE using synthetic oligonucleotides. These oligonucleotides contain nucleotide sequences flanking the desired mutation sites. Mutations may be generated in the full-length DNA sequence of ICE (p45) or in any sequence coding for p30, or p20 or p10 polypeptides.

According to this invention, a mutated ICE
DNA sequence produced by the methods described above,
or any alternative methods known in the art, can be
expressed using an expression vector. An expression
25 vector, as is well known in the art, typically includes
elements that permit autonomous replication in a host
cell independent of the host genome, and one or more
phenotypic markers for selection purposes. Either
prior to or after insertion of the DNA sequences
30 surrounding the desired ICE mutant coding sequence, an
expression vector also will include control sequences
encoding a promoter, operator, ribosome binding site,
translation initiation signal, and, optionally, a
repressor gene or various activator genes and a signal
35 for termination. In some embodiments, where secretion

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of the produced mutant is desired, nucleotides encoding a "signal sequence" may be inserted prior to the ICE mutant coding sequence. For expression under the direction of the control sequences, a desired DNA sequence must be operatively linked to the control sequences -- i.e., they must have an appropriate start signal in front of the DNA sequence encoding the ICE mutant and maintaining the correct reading frame to permit expression of that sequence under the control of the control sequences and production of the desired product encoded by that ICE sequence.

Any of a wide variety of well known available expression vectors are useful to express the mutated ICE coding sequences of this invention.

These include, for example, vectors 15 consisting of segments of chromosomal, non-chromosomal and synthetic DNA sequences, such as various known derivatives of SV40, known bacterial plasmids, e.g., plasmids from E. coli including col E1, pCR1, pBR322, 20 pMB9 and their derivatives, wider host range plasmids, e.g., RP4, phage DNAs, e.g., the numerous derivatives of phage λ , e.g., NM 989, and other DNA phages, e.g., M13 and filamentous single stranded DNA phages, yeast plasmids such as the 2μ plasmid or derivatives thereof, 25 and vectors derived from combinations of plasmids and phage DNAs, such as plasmids which have been modified to employ phage DNA or other expression control sequences. In the preferred embodiments of this invention, we employ E. coli vectors.

In addition, any of a wide variety of expression control sequences -- sequences that control the expression of a DNA sequence when operatively linked to it -- may be used in these vectors to express the mutated DNA sequences according to this invention. Such useful expression control sequences, include, for

- 29 -

example, the early and late promoters of SV40 for animal cells, the lac system, the trp system the TAC or TRC system, the major operator and promoter regions of phage λ the control regions of fd coat protein, all for E. coli, the promoter for 3-phosphoglycerate kinase or other glycolytic enzymes, the promoters of acid phosphatase, e.g., Pho5, the promoters of the yeast α-mating factors for yeast, and other sequences known to control the expression of genes of prokaryotic or eukaryotic cells or their viruses, and various combinations thereof. In the preferred embodiments of this invention, we employ either E. coli or eukaryotic expression in COS-1 cells, a monkey kidney cell line.

A wide variety of hosts are also useful for producing mutated ICE according to this invention. These hosts include, for example, bacteria, such as E. coli, Bacillus and Streptomyces, fungi, such as yeasts, and animal cells, such as CHO and COS-1 cells, plant cells and transgenic host cells. In preferred embodiments of this invention, the host cells are E. coli or COS-1 cells.

It should be understood that not all expression vectors and expression systems function in the same way to express mutated DNA sequences of this invention and to produce modified ICE or ICE mutants. Neither do all hosts function equally well with the same expression system. However, one of skill in the art may make a selection among these vectors, expression control sequences and hosts without undue experimentation and without departing from the scope of this invention. For example, an important consideration in selecting a vector, will be the ability of the vector to replicate in a given host. The copy number of the vector, the ability to control that copy number, and the expression of any other

proteins encoded by the vector, such as antibiotic markers, should also be considered.

In selecting an expression control sequence, a variety of factors should also be considered. These include, for example, the relative strength of the system, its controllability, its compatibility with the DNA sequence encoding the modified ICE of this invention, particularly with regard to potential secondary structures.

Hosts should be selected by consideration of their compatibility with the chosen vector, the toxicity of the modified ICE to them, their ability to secrete mature products, their ability to fold proteins correctly, and to form tetramers, their fermentation requirements, the ease of the purification of the modified ICE from them and safety. Within these parameters, one of skill in the art may select various vector/expression control system/host combinations that will produce useful amounts of the mutant ICE.

The mutant ICE produced in these systems may be purified by a variety of conventional steps and strategies, including those used to purify wild-type ICE.

Once the ICE mutants have been generated in the desired location, i.e., active site or accessory binding site, the mutants may be tested for any one of several properties of interest.

altered charge at physiological pH. This is determined by measuring the mutant ICE isoelectric point (pI) in comparison with that of the wild-type parent.

Isoelectric point may be measured by gelelectrophoresis according to the method of Wellner, D., Analyt. Chem., 43, p. 597 (1971). A mutant with an altered surface charge is an ICE polypeptide containing

- 31 -

a replacement amino acid located at the surface of the enzyme, as provided by the structural information of this invention, and an altered pI.

Furthermore, mutants may be screened for high specific activity in relation to the wild-type ICE. A mutant would demonstrate high specific activity if its second order rate constant (K_{cat}/K_m) for hydrolysis of the substrate Ac-Tyr-Val-Ala-Asp-amino methylcoumarin exceeds 7 x 10⁴ $\rm K^{-1}s^{-1}$ at 25°C, using the assay in

10 Pennington & Thornberry, supra.

A mutant would be tested for altered ICE substrate specificity by measuring the hydrolysis of fluorgenic peptide substrates or unmodified ICE peptide substrates as described in Thornberry et al., supra.

- An enzyme with altered substrate specificity is an enzyme whose second order rate constant (k_{cat}/K_m) for a substrate X_1 -Tyr-Val-Ala- X_2 - X_3 that exceeds the k_{cat}/K_m for the analogous peptide substrate X_1 -Tyr-Val-Ala-Asp- X_3 . X_1 is an amino protecting group, such as
- acetyl; X_2 is a natural or unnatural amino acid residue other than L-aspartate; X_3 is a carboxyl protecting group such as aminomethylcoumarin or p-nitroaniline.

Further properties of interest also include mutants with increased stability to subunit

- 25 dissociation. An ICE mutant with increased stability to subunit dissociation would demonstrate no loss of enzymic activity at concentrations of the enzyme below 10 nM in comparison with the wild-type ICE, which demonstrates a Kd between 1-10 nM.
- In order that the invention described herein may be more fully understood, the following examples are set forth. It should be understood that these examples are for illustrative purposes only and are not to be construed as limiting this invention in any manner.

EXAMPLE 1

Crystal Structure of ICE

The cDNA encoding the p30 precursor of active human ICE (residues Asn 120 to His 404 of p45

(Thornberry et al., <u>supra</u>) was cloned into a P_L promoter expression vector (provided by Dr. J. Mankovich) and expressed in <u>E. coli</u> by temperature-shift induction.

Pre-induction repression of the P_L promoter

10 was achieved by co-expression of the c<u>I</u> repressor gene
on a co-resident, compatible plasmid (pACYC184cI) in
the <u>E. coli</u> host, JM109. Yanish-Perron, C., et al.,
Gene, 33, pp. 103-199 (1985; ATCC #53323). The
promoter was induced by increasing the temperature from
15 28°C to 42°C, at which point the temperature sensitive
c<u>I</u> repressor gene product denatures and gene expression
is initiated, directed by the P_L promoter. Maintenance
of the temperature at 42°C for a further 4 hours
resulted in the accumulation of high levels of the
inactive ICE p30 precursor product within the host cell
cytoplasm, in the form of inactive inclusion bodies.

After mechanical disruption of the cells, and harvesting of the insoluble fraction, the inclusion bodies were washed by suspension in 2M urea, 25mM tris, 0.5mM DTT, 0.1mM EDTA, 0.1mM PMSF, pH 7.5 at 4°C, followed by centrifugation. The inclusion bodies were solubilized in the above buffer containing 7M urea, centrifuged and subjected to size-exclusion chromatography in the same buffer. The p30 fractions, identified by SDS-PAGE and N-terminal sequence analysis, were pooled and diluted to 0.3 mg/ml using column buffer. This was followed by dialysis at 4°C against 25mM tris, 1mm DTT, pH 7.5, until the urea

- 33 -

concentration was less than 20mM, thereby allowing the enzyme to refold.

The protein was concentrated to 3-5 mg/ml by ultrafiltration at 4°C, followed by incubation at room temperature. The disappearance of the p30 precursor, and the concomitant appearance of the p20 and p10 subunits, was monitored by SDS-PAGE, evidence that autocatalytic processing of the enzyme had occurred. ICE enzymatic activity was assayed by hydrolysis of a Succinyl-Tyr-Val-Ala-Asp-p-nitroanilide substrate at 37°C and correlated closely with conversion to subunits.

The autoprocessed ICE was inhibited fully by adding a 2x molar excess of a tetrapeptide aldehyde

15 inhibitor (acetyl-Tyr-Val-Ala-Asp-H). The protein-inhibitor complex sample was concentrated and fractionated by size-exclusion chromatography, in final preparation for crystallization experiments.

Crystals of ICE in complex with the inhibitor

20 were grown by vapor diffusion. Davies, D.R. and D.M.

Segal, Meth. Enzymol., 22, p. 266 (1971). Protein (20

mg/ml in 50 mM citrate, 2.0 mM DTT, pH 6.5) was mixed

with an equal volume of reservoir buffer (15% (w/v) PEG

4K, 400 mM LiSO₄, 200 mM sodium Hepes, 5mM sodium

25 cacodylate, 0.5% beta-octyl glucoside, pH 7.0) and

allowed to stand over the reservoir solution at 4°C.

Crystals grew over a six week period to form tetragonal

bipyramids and were equilibrated with 18% PEG 4K,

400 mM LiSO₄, 200 mM sodium Hepes, 5mM sodium

30 cacodylate, 0.5% beta-octyl glucoside, pH 7.0 prior to

data collection or heavy atom derivatization.

Those of skill in the art will appreciate that the aforesaid crystallization conditions can be varied. Such variations may be used alone or in combination, and include final protein/inhibitor

complex concentrations between 5 mg/ml and 35 mg/ml; all combinations of ICE/inhibitor to precipitant ratios; citrate concentrations between 1mM and 200 mM; DTT concentrations between 0 mM and 10 mM; and any concentration of ß-mercaptoethanol; pH ranges between 5.5 and 9.5; PEG concentrations between 10% and 25% (gms/100ml); PEG weights between 2000 and 8000; LisO4 concentrations between 50 and 750 mM; HEPES concentrations between 5 and 395 mM; and any concentration or type of detergent; any temperature between -5°C and 30°C; and crystallization of ICE/inhibitor complexes by batch, liquid bridge, or dialysis method using these conditions or variations thereof.

R-axis IIC image plate system except for the 2.2Å

Synchrotron data set that was used for refinement of the final model. This data was collected at Cornell

High Energy Synchrotron Source ("CHESS") on a charge
couple device and was reduced to structure factor amplitudes using the Denzo Software Package (Denzo - An Oscillation Data Processing Program For Macro Molecular Crystallography, ©1993, Daniel Gewirth, Yale

University). Oscillation photographs were integrated and reduced to structure factor amplitudes using software supplied by the manufacturer (Molecular Structures Corp., Dallas, Texas).

Refined heavy atom parameters were used to compute multiple isomorphous replacement phases.

Inclusion of the anomalous data for the Hg derivative in cross-phased difference Fourier maps showed the space group to be P4₃2₁2 rather than its enantiomorph. The mean figure of merit, including anomalous data for the Hg derivative, was 0.65 to 3.5Å resolution (Table 35 1).

Solvent flattening and phase extension (CCP4-Collaborative Computing Project No. 4, A Suite of Programs for Protein Crystallography; Daresbury Laboratory, Warrington, WA4 4AD, U.K. (1979)) improved the map and allowed identification of some of the residues in the protein core. Cycles of model building (Quanta, version 4.0b, Molecular Simulations Inc., Burlington MA), positional refinement, (Brunger, A.T., J. Acta Cryst., A46, pp. 46-57 (1990); Brunger, A.T. et al., J. Acta Cryst., A46, pp. 585-93 (1990)) and phase combination (CCP4-Collaborative Computing

Project, <u>supra</u>) were carried out until the switch to phases calculated from the model could be made.

Refinement continued against the -16°C, 2.2Å data

15 (Table 1), which allowed the more difficult loop regions of the protein to be constructed.

The following table summarizes the X-ray crystallography data sets of ICE derivatives that were used to determine the structure of ICE according to 20 this invention.

Table 1

	Protein Modification	Resolution A	Complete- ness of data %	Rmerge %		t ceil sions, Å c	No. of	Rc %	Phasing Power
25	Tetrapeptide aldehyde*	20 - 2.2	87	7.1	64.9	164.1			-
	Tetrapeptide aldehyde * *	20 - 2.6	90	8.3	64.4	163.3	••	_	
	Tetrapeptide aldehyde	20 - 2.8	78	8.3	64.7	162.9	-	_	••
	kodinated tetrapeptide aldehyde	20 - 3.5	86	9.4	64.4	162.8	2	0.88	1.09
30	Thimerosal	20 - 3,5	88	8.4	64.4	162.3	5	0.67	1.08
	Gold Thiomalats	20 - 3.5 .	74	9.5	64.7	162.7	3	0.72	1,22
	Uranyl Acetate	20 - 4.0	80	10.8	64.7	162.9	2	0.79	1.32
	Lead Chloride	20 - 3.5	64	8.9	64.7	162.8	2	0.76	1.38
	* Data collected at -16°C	at CHESS			• • • • • • • • • • • • • • • • • • • •	.02.5	•	0.70	1.30
35	** Data collected at -16°	C							

<u>Definitions</u>: Rmerge gives the agreement between repeated intensity measurements, with the number of crystals used in the data set given in parentheses. The number of heavy-atom binding sites is given where

appropriate. R_c is the Cullis R factor for centrosymmetric reflections, and the phasing power is the ratio of average heavy-atom scattering to the average lack of closure of the phase triangles.

5 Blundell, T.L. and Johnson, L.N., <u>Protein</u> <u>Crystallography</u>, Academic Press, New York (1976).

The ICE tetrameric model according to this invention has an R-factor of 19% against all observed data between 7Å and 2.2Å resolution, with

10 root-mean-square deviation from ideal bond lengths and angles of 0.011Å and 2.84Å respectively.

EXAMPLE 2

Confirmation of the Active Site of ICE

In order to confirm the location of the

15 active site in the tetrameric ICE molecule, as deduced
from the structure coordinates of ICE, a series of p30
ICE mutants was generated.

Oligonucleotide-directed mutagenesis was
performed on pcDNA3 (Invitrogen) constructs using
uracil-enrichment of single-strand DNA. Kunkel, T.A.,
<u>Proc. Nat. Acad. Sci.</u>, 82, pp. 488-492 (1985); Kunkel,
T.A. et al., <u>Meth. Enzymol.</u>, 154, pp. 367-382 (1987).
This is a modification of the method originally
described for M13 mutagenesis. Zoller, M.J. and
M. Smith, <u>Nucleic Acids Res.</u>, 10, pp. 6487-6500 (1983);
Zoller M.J. and M. Smith, <u>Meth. Enzymol.</u>, 100, pp. 468-

Mutagenesis was performed using the reagents provided in the Muta-Gene Kit (BioRad). Mutagenesis primers were synthesized in the (+) coding orientation. The <u>dutung E. coli</u> strain CJ326 was used for uracil enrichment of single-strand DNA, and the MV1190 strain was used for selection of heteroduplex DNA after extension-ligation reactions. All oligonucleotides

- 37 -

were synthesized on an applied Biosystems 380 DNA synthesizer and purified by electrophoreses in polyacrylamide-urea slab gels. Mutations made in the 30kDa ICE-encoding cDNA were fully sequenced in the coding region by the dideoxy method. Sanger, F. et al., Proc. Nat. Acad. Sci. 74, pp. 5463-5467 (1977). Mutant DNA in preparation for COS-1 cell transfection, or alternatively E. coli transfection, was purified by alkaline lysis and cesium gradient centrifugation prior to transfection.

Each mutant cDNA was transfected into a COS1 cell line, then tested for its ability to process
pIL-1ß in vitro, i.e., to secrete mature IL-1ß. The
COS-1 cell line used, had previously been transfected
with a pIL-1ß encoding cDNA cloned into an MNC stuffer
vector (B. Seed, Harvard Medical School) which had
subsequently integrated into the chromosome. pIL-1ß
production was maintained by the addition of 0.5 mg/ml
G-418 Sulfate to culture media.

Approximately 3 \times 10⁶ COS-1 cells in 100mm 20 tissue culture plates were transfected with 15 $\mu\mathrm{g}$ of each plasmid. DNA was mixed with 200 μ l DEAE-Dextran, brought to 4 ml with phosphate-buffered saline, and added to the plates. Cells were incubated at 37°C for 25 30 min. 8 ml of an 80 μ M chloroquine/serum-free DMEM solution was added and the cells were incubated for 2.5 hr. This solution was aspirated and cells were treated for two minutes with 10% DMSO/serum-free DMEM. After washing with serum-free media, 10 ml complete 30 media was added. Conditioned media were sampled at 16 and 24 hr. Activity in this assay requires that transcription, translation and protein folding of mutants are not arrested. The amount of mutant ICE present in cell lysates was determined by Western blot 35 using an anti-p20 rabbit antiserum that recognizes

amino acids 136-150 inclusive and which also recognizes the intact p30 precursor.

Mature IL-1ß in the cell medium was detected by ELISA (R&D Systems). Samples were diluted to achieve concentrations in the linear range of the ELISA assay (8-60 pg/ml). Background IL-1ß levels were determined in cells transfected with the expression vector lacking ICE cDNA, and this value was subtracted from all other concentrations. The % activity values were calculated as the ratio of secreted IL-1ß from cells transfected with mutant ICE divided by IL-1ß secreted by cells transfected with wild-type ICE. The final ratio is the mean of at least two experiments. These data are recorded in Figure 3.

Based on these data, it was determined that
mutation of Cys-285 or His-237 eliminates pIL-16
processing activity, as well as autoprocessing.
Mutation of Arg-179, which contacts the Pl Asp to Glu,
also abolishes activity. Mutation of Cys-244 to Ala,
which may contact P' side chains of substrates, reduces
enzymatic activity significantly. In contrast,
mutation of other residues proximal to Cys-285
including Ser-332, -333 or -339, and His-249, does not
eliminate activity. Accordingly, we confirmed the
importance of various residues in the ICE active site.

EXAMPLE 3

The Use of Molecular Replacement To Solve An Unknown ICE Crystal Structure

The method of molecular replacement was used to determine the structure coordinates of crystals of ICE in complex with the tetrapeptide aldehyde inhibitor Ac-Tyr-Val-Pro-Asp-H in comparison with crystals of ICE in complex with the tetrapeptide aldehyde inhibitor Ac-Tyr-Val-Ala-Asp-H (as prepared in Example 1).

- 39 -

Crystals of ICE in complex with the tetrapeptide aldehyde inhibitor, Ac-Tyr-Val-Pro-Asp-H ("Pro") were grown under conditions identical to those for crystals of ICE in complex with the tetrapeptide aldehyde inhibitor, Ac-Tyr-Val-Ala-Asp-H ("Ala").

X-ray diffraction data to 2.8Å resolution was collected on the ICE/Pro co-complex. A difference electron density map that combined diffraction data of the form | F_{Pro} - F_{Ala} | and phases calculated from the refined model of the Ala inhibited enzyme was used to locate structure changes that had occurred in the ICE/Pro co-complex.

Negative features were found in the map wherever localized atoms in the Ala complex were

15 removed or shifted by switching to the new ligand.

Positive features were found when localized atoms were introduced into the structure, and indicated the new positions of shifted atoms.

Replacement of the alanine that sits in the

P2 binding pocket in Ala with proline in Pro introduced
two methylene groups into the structure of the ICE cocomplex. The location of these new atoms was indicated
by the presence of positive difference electron density
adjacent to the beta-carbon of the alanine in the

binding pocket P2. Another positive peak nearby
indicated the binding of a new water molecule in the
Pro complex relative to the Ala complex. There were
also pairs of positive and negative peaks near the
tyrosine moiety that sits in the P4 binding pocket of
the inhibitor. These peaks indicated shifts in the
position of these atoms in the Pro complex relative to
their location in the Ala complex.

These shifts, plus the new atoms referred to above, were modeled, and the resulting structure was refined against the X-ray data to determine a final

- 40 -

picture of the co-complex of Pro with ICE. The space group $(P4_32_12)$ and unit cell dimensions $(a=b=65\pm 5 \text{\AA})$ c=162 \pm 5 \AA) for the Pro complex were the same as those observed for the Ala complex.

The ICE structure coordinates known for the first time by virtue of this invention may be used to solve the unknown structure of any mutant, homologue or co-complex of ICE using the above-described method.

This method may also be used to determine the binding or orientation of a ligand or chemical entity in the active site or accessory binding site of ICE.

While we have described a number of embodiments of this invention, it is apparent that our basic examples may be altered to provide other embodiments which utilize the products and processes of this invention. Therefore, it will be appreciated that the scope of this invention is to be defined by the appended claims rather than by the specific embodiments which have been represented by way of example.

Tables A and B, following this page, list respectively, the tetramer interface contacts and the structure coordinates of the ICE molecule of this invention.

20

- 41 -

TABLE A

TETRAMER INTERFACE CONTACTS

	Re	sidue	R	esidue	R	esi	.due	
	<u>P20</u>	<u>P10</u> .	<u>P10</u>	<u>P10</u>	<u>P</u>	20	<u>P20</u>	
5	150		. 320	382	. 2	40	259	
	151	371	320	380	2	67	293	
	151	372	322	377	2	68	293	
		375	322	378	2	74	295	
	291	323	322	380				
10	291	321	322	384				
•	291	322	322	385				
	291	323	322	386				
	292	321	323	327				
	293	321	324	334				
15	293		324	386				
	293	320	325	378				
	294	318	. 325	386				
	294		334	393	•			
	294			391				
20			367	367				
	295		367	374				
		320	371	394				
		321	371	395				
		317	371	396				
25	297	317	374					
			374	393				
			374					
				395				
				396				
30			•	395				
				396				
			386					
				395				
				392				
35			388					
			388					
			389					
			389					
			390	391				

- 42 -

TABLE B

STRUCTURE COORDINATES OF ICE

ATOM				Atom Type	RESIDUE		<u>x</u>	<u>_Y_</u>		<u>occ</u>	<u>B</u>
ATOM 2 C GLY 131 50.203 22.856 9.789 1.00 67.00 ATOM 3 HT1 GLY 131 51.316 82.335 -11.694 1.00 0.00 ATOM 4 HT2 GLY 131 50.746 82.335 -11.694 1.00 0.00 ATOM 5 N GLY 131 50.746 82.335 -11.694 1.00 0.00 ATOM 6 HT3 GLY 131 50.746 82.310 -12.180 1.00 0.00 ATOM 6 HT3 GLY 131 50.783 81.456 -13.079 1.00 0.00 ATOM 7 CA GLY 131 50.783 81.456 -13.079 1.00 0.00 ATOM 8 N ASN 132 49.175 80.9016 8.934 1.00 65.90 ATOM 9 H ASN 132 49.175 81.466 -7.566 1.00 57.33 ATOM 10 CA ASN 132 49.178 81.466 -7.566 1.00 57.33 ATOM 11 CB ASN 132 47.578 81.260 -6.979 1.00 65.76 ATOM 12 CG ASN 132 47.590 82.132 5.758 1.00 65.77 ATOM 13 ODI ASN 132 44.506 81.800 4.977 1.00 65.77 ATOM 14 ND2 ASN 132 44.506 81.800 4.977 1.00 65.71 ATOM 15 ED21 ASN 132 44.506 81.800 4.977 1.00 67.11 ATOM 16 HD22 ASN 132 45.599 81.00 5.228 1.00 0.00 ATOM 16 HD22 ASN 132 45.999 81.00 5.228 1.00 0.00 ATOM 16 HD22 ASN 132 50.481 80.946 5.521 1.00 0.00 ATOM 17 C ASN 132 50.481 80.946 5.521 1.00 0.00 ATOM 18 O ASN 132 50.481 80.946 5.521 1.00 0.00 ATOM 19 N VAL 133 50.767 79.856 6.277 1.00 67.11 ATOM 19 N VAL 133 50.767 79.856 6.277 1.00 67.11 ATOM 22 CB VAL 133 50.767 79.856 6.277 1.00 67.11 ATOM 22 CG VAL 133 51.801 79.99 81.00 5.577 1.00 67.11 ATOM 24 CC2 VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 25 CG VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 26 CD VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 27 CG VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 28 NZ LYS 134 53.901 79.99 81.00 6.377 1.00 67.11 ATOM 29 CA VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 20 CG VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 27 CG VAL 133 51.801 79.99 81.00 6.377 1.00 67.11 ATOM 28 NZ LYS 134 53.901 79.99 81.00 6.370 1.00 67.11 ATOM 30 CE LYS 134 53.901 79.99 81.00 6.370 1.00 67.11 ATOM 30 CE LYS 134 53.90 80.901 7.00 1.00 67.11 ATOM 30 CE LYS 134 53.90 80.901 7.00 1.00 67.11 ATOM 30 CE LYS 134 53.90 80.901 7.00 1.00 67.00 67.90 ATOM 40 NZ LYS 134 53.90 80.90 80.903 1.10 67.90 ATOM 40 NZ LYS	_			•	CIV	121	AD 9.49	81 525	-9.909	1.00	66.26
ATOM 3 HTI GLY 131 51.316 82.385 -11.694 1.00 0.00 ATOM 4 HTZ GLY 131 50.346 82.510 -12.180 1.00 0.00 ATOM 6 HT3 GLY 131 50.746 82.510 -12.180 1.00 0.00 ATOM 6 HT3 GLY 131 50.783 81.456 -13.079 1.00 0.00 ATOM 7 CA GLY 131 50.783 81.456 -13.079 1.00 65.90 ATOM 8 N ASN 132 49.175 80.916 -8.934 1.00 65.90 ATOM 9 H ASN 132 49.175 80.916 -8.934 1.00 65.90 ATOM 10 CA ASN 132 49.178 81.466 -7.566 1.00 0.00 ATOM 11 CB ASN 132 47.778 81.466 -7.566 1.00 53.76 ATOM 12 CG ASN 132 47.550 82.132 -5.758 1.00 65.71 ATOM 13 OD1 ASN 132 44.506 81.800 -4.997 1.00 65.21 ATOM 14 ND2 ASN 132 45.919 81.109 -5.228 1.00 65.21 ATOM 15 HD21 ASN 132 45.919 81.109 -5.228 1.00 0.00 ATOM 16 HD22 ASN 132 50.261 80.777 -6.766 1.00 57.45 ATOM 18 O ASN 132 50.467 80.946 -4.223 1.00 0.00 ATOM 19 N VAL 133 50.972 79.911 -7.425 1.00 46.19 ATOM 19 N VAL 133 50.972 79.911 -7.425 1.00 46.19 ATOM 22 CB VAL 133 53.081 79.994 -6.973 1.00 47.07 ATOM 24 CG2 VAL 133 53.081 79.994 -6.973 1.00 42.45 ATOM 25 CVAL 133 53.5076 79.985 -7.099 1.00 42.45 ATOM 26 CVAL 133 53.5081 79.994 -6.973 1.00 42.45 ATOM 27 N LYS 134 53.91 79.997 -5.986 1.00 35.70 ATOM 28 NC VAL 133 53.081 79.994 -6.973 1.00 42.45 ATOM 29 CA LYS 134 53.91 79.994 -5.226 1.00 47.07 ATOM 20 CH VAL 133 50.765 79.856 -7.099 1.00 42.45 ATOM 21 CA VAL 133 53.081 79.994 -6.973 1.00 42.45 ATOM 22 CB VAL 133 53.081 79.994 -6.973 1.00 42.45 ATOM 24 CG2 VAL 133 53.511 88.670 -8.000 1.00 43.94 ATOM 27 N LYS 134 55.912 79.956 -7.099 1.00 42.45 ATOM 27 N LYS 134 55.912 79.956 -7.099 1.00 42.55 ATOM 30 CB LYS 134 55.908 81.371 -7.425 1.00 47.07 ATOM 30 CB LYS 134 55.908 81.371 -7.425 1.00 47.07 ATOM 30 CB LYS 134 55.908 81.371 -7.426 1.00 5.37 ATOM 31 CG LYS 134 57.679 81.391 -7.425 1.00 40.99 ATOM 30 CB LYS 134 55.606 79.261 -7.445 1.00 5.99 ATOM 40 N LEU 135 56.807 82.261 -7.445 1.00 5.99 ATOM 40 N LEU 135 56.807 82.261 -7.445 1.00 5.99 ATOM 40 N LEU 135 56.807 82.261 -7.445 1.00 5.99 ATOM 40 CB LYS 134 57.699 81.591 -7.426 1.00 5.99 ATOM 40 CB LYS 134 55.606 79.261 -7.445 1.00 5.99 ATOM 40 CB LEU 135 56.807	5									1.00	67.80
ATOM 4 HTZ GLY 131 49.746 82.510 -12.180 1.00 0.00 ATOM 5 N GLY 131 50.546 81.841 -12.188 1.00 17.76 ATOM 6 HT3 GLY 131 50.783 81.456 13.079 1.00 0.00 ATOM 7 CA GLY 131 50.783 81.456 13.079 1.00 65.50 ATOM 8 N ASN 132 49.175 80.916 8.934 11.00 63.34 ATOM 10 CA ASN 132 49.175 80.916 8.934 11.00 63.34 ATOM 11 CB ASN 132 47.778 81.260 6.979 1.00 65.76 ATOM 11 CB ASN 132 47.778 81.260 6.979 1.00 65.77 ATOM 13 ODI ASN 132 47.590 82.132 5.758 1.00 65.77 ATOM 14 ND2 ASN 132 47.590 82.132 5.758 1.00 65.77 ATOM 15 ND2 ASN 132 47.590 82.132 5.758 1.00 65.77 ATOM 16 ND2 ASN 132 45.999 81.100 5.760 6.711 ATOM 17 C ASN 132 45.999 81.100 5.228 1.00 0.00 ATOM 16 HD22 ASN 132 45.999 81.100 5.228 1.00 0.00 ATOM 17 C ASN 132 50.487 80.946 5.521 1.00 0.00 ATOM 18 N VAL 133 50.765 79.836 8.377 6.766 1.00 51.45 ATOM 19 N VAL 133 50.765 79.836 8.377 1.725 1.00 0.00 ATOM 19 N VAL 133 50.765 79.836 8.377 1.725 1.00 0.00 ATOM 22 CB VAL 133 50.787 79.991 1.00 42.45 ATOM 22 CG VAL 133 51.081 79.994 5.711 1.00 39.72 ATOM 22 CB VAL 133 51.081 79.994 5.711 1.00 39.72 ATOM 23 CG1 VAL 133 50.765 79.836 8.377 1.00 42.45 ATOM 24 CG2 VAL 133 51.511 80.670 8.070 1.00 42.45 ATOM 25 C VAL 133 51.511 80.670 8.000 1.00 42.45 ATOM 26 O VAL 133 51.511 80.670 8.000 1.00 42.45 ATOM 27 N LYS 134 55.201 81.37 4.424 1.00 39.72 ATOM 28 C VAL 133 51.511 80.670 8.000 1.00 42.45 ATOM 38 NZ LYS 134 55.201 81.37 4.424 1.00 58.49 ATOM 39 CB LYS 134 55.802 80.804 4.488 1.00 47.93 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.801 81.37 4.224 1.00 0.00 4.00 ATOM 30 CB LYS 134 55.800 80.873 1.10 0.40 4.00 4.00 ATOM 30 CB LYS 134 55.800 80.873 1.10 0.40 4.00 4.00 ATOM 30 CB LYS 134 55.800 80.873 1.10 0.40 4.00 4.00 ATOM 40 CB LEU 135 56.800 8				_						1.00	0.00
ATOM 5 NN GLY 131 50.546 81.841 -12.148 1.00 71.76 ATOM 6 HTJ GLY 131 50.546 81.841 -12.148 1.00 71.76 ATOM 7 CA GLY 131 50.783 81.456 -13.079 1.00 65.90 ATOM 8 N ASN 132 49.175 80.916 -8.934 1.00 65.90 ATOM 9 H ASN 132 49.175 80.916 -8.934 1.00 65.90 ATOM 10 CA ASN 132 49.178 81.466 -7.566 1.00 57.35 ATOM 11 CB ASN 132 47.778 81.260 6-979 1.00 65.76 ATOM 12 CG ASN 132 47.759 81.260 6-979 1.00 65.76 ATOM 14 ND2 ASN 132 47.550 82.132 -5.738 1.00 65.77 ATOM 15 ODI ASN 132 45.950 82.132 -5.738 1.00 65.71 ATOM 16 HD22 ASN 132 45.950 81.860 4.997 1.00 65.21 ATOM 15 HD21 ASN 132 46.362 82.459 -5.228 1.00 0.00 ATOM 16 HD22 ASN 132 46.382 82.459 -5.228 1.00 0.00 ATOM 16 HD22 ASN 132 50.261 80.717 -6.706 1.00 51.45 ATOM 19 N VAL 133 50.767 79.951 1.00 46.19 ATOM 20 H VAL 133 50.767 79.951 1.00 46.19 ATOM 21 CA VAL 133 50.767 79.856 -8.373 1.00 0.00 ATOM 22 CB VAL 133 50.261 80.777 -8.956 -8.373 1.00 0.00 ATOM 23 CG1 VAL 133 50.888 77.172 -8.00 1.00 35.75 ATOM 24 CG2 VAL 133 50.888 77.172 -8.00 1.00 35.70 ATOM 25 C VAL 133 50.868 77.172 -8.00 1.00 35.70 ATOM 26 O VAL 133 50.868 77.172 -8.00 1.00 35.70 ATOM 27 N LYS 134 55.42 80.670 -8.000 1.00 43.94 ATOM 29 CA LYS 134 55.942 80.670 -8.000 1.00 43.94 ATOM 30 CB LYS 134 57.202 80.832 -7.099 1.00 42.39 ATOM 20 CA LYS 134 57.202 80.832 -7.099 1.00 42.39 ATOM 20 CD LYS 134 57.603 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.603 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.603 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.603 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.508 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.509 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.509 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.509 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.509 81.341 -7.997 1.00 68.37 ATOM 30 CB LYS 134 57.509 81.341 -7.997 1.00 68.37 ATOM 40 N LEU 135 56.807 82.320 -7.039 1.00 40.994 ATOM 40 N LEU 135 56.807 82.320 -7.039 1.00 40.995 ATOM 40 CD LYS 134 57.509 81.341 -7.997 1.00 68.37 ATOM 40 N LEU 135 56.807 82.320 -7.035 1.00 60.00 ATOM 40 CD LYS 134 57.					GLY					1.00	
10 ATOM 6 HTG GLY 131 50.783 81.456 -13.079 1.00 0.00 ATOM 7 CA GLY 131 50.783 81.456 -13.079 1.00 65.90 ATOM 8 N ASN 132 49.175 80.916 -8.934 1.00 65.90 ATOM 10 CA ASN 132 49.175 80.916 -8.934 1.00 6.34 ATOM 9 H ASN 132 49.175 80.916 -8.934 1.00 6.30 ATOM 10 CA ASN 132 49.175 80.916 -8.934 1.00 6.30 6.76 ATOM 11 CB ASN 132 47.778 81.260 -6.979 1.00 65.76 ATOM 11 CB ASN 132 47.778 81.260 -6.979 1.00 65.76 ATOM 12 CG ASN 132 47.590 82.132 -5.758 1.00 65.76 ATOM 13 ODI ASN 132 48.282 83.107 -5.487 1.00 65.71 ATOM 14 ND2 ASN 132 45.919 81.109 -5.228 1.00 0.00 ATOM 16 HD22 ASN 132 45.919 81.109 -5.228 1.00 0.00 ATOM 16 HD22 ASN 132 45.939 81.109 -5.228 1.00 0.00 ATOM 16 HD22 ASN 132 45.939 81.09 -5.228 1.00 0.00 ATOM 18 O ASN 132 50.261 80.777 -6.706 1.00 51.45 ATOM 19 N VAL 133 50.261 80.777 -7.425 1.00 47.07 ATOM 19 N VAL 133 50.972 79.911 -7.425 1.00 47.07 ATOM 20 H VAL 133 50.972 79.911 -7.425 1.00 47.07 ATOM 20 H VAL 133 50.972 79.911 -7.425 1.00 42.45 ATOM 22 CB VAL 133 53.081 79.994 6-9.973 1.00 42.45 ATOM 22 CB VAL 133 53.081 79.994 6-9.973 1.00 42.45 ATOM 22 CB VAL 133 53.081 79.994 6-9.973 1.00 42.45 ATOM 24 CG2 VAL 133 53.302 79.856 8.373 1.00 0.00 ATOM 25 CC VAL 133 53.303 79.995 6-7.538 1.00 39.93 ATOM 24 CG2 VAL 133 53.303 79.995 6-7.538 1.00 39.93 ATOM 24 CG2 VAL 133 53.303 79.995 6-7.538 1.00 39.93 ATOM 25 CC VAL 133 53.301 79.997 6-9.986 1.00 42.49 ATOM 29 CA LYS 134 54.119 79.997 5-9.986 1.00 43.43 ATOM 20 CD LYS 134 54.119 79.997 5-9.986 1.00 43.43 ATOM 20 CD LYS 134 55.401 80.832 2.2773 1.00 68.37 ATOM 24 CG2 VAL 133 53.301 80.832 -7.993 1.00 40.00 ATOM 24 CG2 VAL 133 53.301 80.832 -7.993 1.00 40.00 ATOM 24 CG2 VAL 133 53.301 80.832 -7.993 1.00 40.00 ATOM 24 CG2 VAL 133 53.301 80.832 -7.993 1.00 40.90 ATOM 25 CC VAL 133 53.301 80.832 -7.993 1.00 40.90 ATOM 25 CC VAL 133 53.301 80.832 -7.993 1.00 40.90 ATOM 25 CC VAL 133 53.301 80.832 -7.993 1.00 40.90 ATOM 25 CC VAL 133 53.301 80.832 -7.993 1.00 40.90 ATOM 25 CC LYS 134 57.90 80.832 -7.993 1.00 40.90 ATOM 25 CC LYS 134 57.90 80.832 -7.993 1.00 4										1.00	71.76
ATOM 7 CA GLY 131 50.192 80.805 -11.207 1.00 65.90 ATOM 8 N ASN 132 49.175 80.805 -11.207 1.00 65.90 ATOM 9 H ASN 132 49.175 80.805 -11.207 1.00 65.90 ATOM 10 CA ASN 132 49.178 81.466 7.566 1.00 57.36 ATOM 11 CB ASN 132 47.778 81.266 -6.799 1.00 65.36 ATOM 12 CG ASN 132 47.550 82.132 -5.738 1.00 65.77 ATOM 13 ODI ASN 132 47.550 82.132 -5.738 1.00 65.77 ATOM 14 ND2 ASN 132 46.506 81.860 4.997 1.00 65.21 ATOM 15 HD21 ASN 132 46.506 81.860 4.997 1.00 65.21 ATOM 16 HD22 ASN 132 46.506 81.860 4.997 1.00 65.21 ATOM 17 C ASN 132 50.261 80.777 6-706 1.00 57.11 ATOM 19 N VAL 133 50.972 79.911 7-7.425 1.00 47.07 ATOM 20 H VAL 133 50.972 79.911 7-7.425 1.00 47.07 ATOM 20 H VAL 133 50.765 79.856 8.373 1.00 40.707 ATOM 22 CB VAL 133 50.868 77.172 8.010 1.00 39.63 ATOM 23 CG1 VAL 133 50.881 79.094 6.973 1.00 39.93 ATOM 24 CG2 VAL 133 50.881 79.994 6.973 1.00 39.93 ATOM 25 C VAL 133 50.868 77.172 8.010 1.00 39.93 ATOM 26 O VAL 133 50.351 79.994 6.973 1.00 39.93 ATOM 27 N LYS 134 53.321 79.994 -6.973 1.00 39.93 ATOM 28 NZ LYS 134 53.321 79.994 -6.973 1.00 47.07 ATOM 30 CB LYS 134 53.321 79.994 -6.973 1.00 47.07 ATOM 29 CA LYS 134 53.301 79.994 -6.973 1.00 47.07 ATOM 30 CB LYS 134 53.301 79.994 -6.973 1.00 39.93 ATOM 26 O VAL 133 50.868 77.172 -8.010 1.00 39.93 ATOM 27 N LYS 134 53.301 79.994 -6.973 1.00 47.92 ATOM 30 CB LYS 134 53.301 80.832 -7.5986 1.00 47.92 ATOM 30 CB LYS 134 53.301 80.832 -7.999 1.00 42.39 ATOM 30 CB LYS 134 53.911 80.670 -8.000 1.00 41.55 ATOM 30 CB LYS 134 53.911 80.670 -8.000 1.00 47.92 ATOM 30 CB LYS 134 53.911 80.670 -8.000 1.00 47.92 ATOM 30 CB LYS 134 53.91 79.997 -7.009 1.00 47.92 ATOM 30 CB LYS 134 53.91 79.997 -7.009 1.00 47.92 ATOM 30 CB LYS 134 53.91 79.997 -7.009 1.00 47.92 ATOM 30 CB LYS 134 53.91 79.997 -7.009 1.00 47.92 ATOM 30 CB LYS 134 53.91 79.997 -7.009 1.00 47.92 ATOM 31 CG LYS 134 53.90 80.809 80.809 1.00 47.92 ATOM 32 CD LYS 134 56.809 80.809 80.809 1.00 50.90 ATOM 40 CD LYS 134 56.809 80.809 80.809 1.00 1.00 50.90 ATOM 40 CD LYS 134 56.809 80.809 80.809 1.00 1.00 50.90 ATOM 40 CD L	10									1.00	0.00
ATOM 8 N ASN 132 49,175 80,916 -8,934 1.00 63.34 ATOM 9 H ASN 132 49,178 81,466 -7,566 1.00 9.00 ATOM 10 CA ASN 132 49,178 81,466 -7,566 1.00 9.73.35 1.00 63,76 ATOM 11 CB ASN 132 47,778 81,260 -6,979 1.00 63,76 ATOM 13 ODI ASN 132 47,758 81,260 -6,979 1.00 65,77 ATOM 13 ODI ASN 132 42,218 83,107 -5,487 1.00 65,21 ATOM 15 HD21 ASN 132 45,218 83,107 -5,487 1.00 65,21 ATOM 16 HD22 ASN 132 45,919 81,109 -5,228 1.00 0.00 ATOM 16 HD22 ASN 132 45,919 81,109 -5,228 1.00 0.00 ATOM 16 HD22 ASN 132 50,487 89,946 -5,221 1.00 0.00 ATOM 17 C ASN 132 50,487 89,946 -5,521 1.00 46,19 ATOM 18 O ASN 132 50,487 89,946 -5,521 1.00 46,19 ATOM 18 O ASN 132 50,487 89,946 -5,521 1.00 46,19 ATOM 19 N VAL 133 50,972 79,951 -7,425 1.00 47,07 ATOM 20 H VAL 133 50,765 79,856 -8,373 1.00 47,07 ATOM 22 CB VAL 133 53,081 79,094 6-9,731 1.00 42,45 ATOM 22 CB VAL 133 53,081 79,094 6-9,731 1.00 42,45 ATOM 24 CG2 VAL 133 53,081 77,172 8,010 1.00 39,72 ATOM 24 CG2 VAL 133 53,342 76,891 -7,947 1.00 39,63 ATOM 25 C VAL 133 53,342 76,891 -7,947 1.00 39,63 ATOM 26 O VAL 133 53,314 57,693 -7,538 1.00 1.00 41,5 ATOM 27 N LYS 134 54,119 79,997 5,986 1.00 41,5 ATOM 27 N LYS 134 54,119 79,997 5,986 1.00 41,5 ATOM 27 N LYS 134 54,119 79,997 5,986 1.00 43,48 ATOM 29 CA LYS 134 55,301 80,832 5,918 1.00 47,93 ATOM 26 C VAL 133 53,341 57,693 -7,598 1.00 47,93 ATOM 27 N LYS 134 54,119 79,997 5,986 1.00 47,93 ATOM 27 N LYS 134 54,119 79,997 5,986 1.00 47,93 ATOM 27 N LYS 134 54,119 79,997 5,986 1.00 47,93 ATOM 28 NZ LYS 134 57,671 81,322 2,777 1.00 68,37 ATOM 30 CB LYS 134 57,671 81,322 2,777 1.00 68,37 ATOM 34 NZ LYS 134 57,671 81,322 2,777 1.00 68,37 ATOM 34 NZ LYS 134 57,693 82,494 1.00 68,39 ATOM 34 NZ LYS 134 57,693 82,494 1.00 68,39 ATOM 35 CE LYS 134 57,693 82,494 1.00 68,39 ATOM 36 CE LYS 134 57,693 82,394 1.00 47,92 1.00 0.00 ATOM 40 N LEU 135 56,807 81,334 -7,698 1.00 47,93 ATOM 34 NZ LYS 134 57,693 82,316 0.0110 1.00 0.00 0.00 ATOM 40 N LEU 135 56,807 81,334 -7,698 1.00 47,95 ATOM 39 O LYS 134 56,809 80,813 1.10 0.00 0.00 0.00 ATOM 40 N LEU 135 56,807	10									1.00	65.90
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ATOM 10 CA ASN 132 49.178 81.466 -7.566 1.00 57.35 ATOM 11 CB ASN 132 47.778 81.260 6.979 1.00 65.76 ATOM 13 OD1 ASN 132 47.578 81.260 6.979 1.00 65.77 ATOM 13 OD1 ASN 132 47.578 81.260 6.979 1.00 65.77 ATOM 14 ND2 ASN 132 45.98 82.132 5.748 7 1.00 65.71 ATOM 15 HD21 ASN 132 45.919 81.109 5.228 1.00 0.00 ATOM 15 HD21 ASN 132 45.919 81.109 5.228 1.00 0.00 ATOM 16 HD22 ASN 132 50.261 80.777 6.706 1.00 0.00 ATOM 17 C ASN 132 50.487 80.946 5.521 1.00 0.00 ATOM 18 O ASN 132 50.487 80.946 5.521 1.00 47.07 ATOM 19 N VAL 133 50.972 79.911 -7.425 1.00 47.07 ATOM 20 H VAL 133 50.765 79.856 8.373 1.00 0.00 ATOM 20 H VAL 133 50.765 79.856 8.373 1.00 42.45 ATOM 22 CB VAL 133 50.867 79.994 6.973 1.00 42.45 ATOM 23 CG1 VAL 133 50.868 77.172 8.010 1.00 1.00 35.70 ATOM 24 CG2 VAL 133 50.868 77.172 8.010 1.00 1.00 35.70 ATOM 25 C VAL 133 50.302 79.996 7.029 1.00 42.93 ATOM 26 O VAL 133 50.302 79.996 7.029 1.00 42.39 ATOM 27 N LYS 134 54.119 79.997 5.596 1.00 43.94 ATOM 28 NZ LYS 134 55.01 80.694 4.498 1.00 43.83 ATOM 29 CA LYS 134 55.01 80.832 5.5918 1.00 47.92 ATOM 29 CA LYS 134 55.801 80.894 4.498 1.00 47.92 ATOM 30 CB LYS 134 55.802 80.694 4.498 1.00 47.92 ATOM 35 CE LYS 134 55.801 80.893 5.5918 1.00 47.93 ATOM 36 CB LYS 134 55.801 80.893 5.5918 1.00 47.93 ATOM 37 HZ3 LYS 134 55.801 80.893 5.5918 1.00 47.93 ATOM 38 C LYS 134 55.801 80.893 6.994 1.00 47.93 ATOM 38 C LYS 134 55.801 80.893 6.994 1.00 47.93 ATOM 38 C LYS 134 55.801 80.893 6.994 1.00 47.93 ATOM 38 C LYS 134 55.801 80.893 6.994 1.00 47.93 ATOM 39 C LYS 134 55.603 80.893 6.994 1.00 47.93 ATOM 39 C LYS 134 55.603 80.893 6.994 1.00 47.93 ATOM 40 N LEU 135 55.807 81.384 7.7698 1.00 47.93 ATOM 40 N LEU 135 55.807 81.384 7.7698 1.00 47.93 ATOM 40 N LEU 135 55.807 81.384 7.7698 1.00 47.93 ATOM 47 C LEU 135 55.807 81.384 7.7698 1.00 47.39 ATOM 48 C LYS 134 55.600 80.819 8.8610 1.00 47.93 ATOM 49 N CYS 136 61.688 80.557 1.00.337 1.00 60.27 ATOM 49 N CYS 136 61.688 80.557 1.00.337 1.00 53.47 ATOM 50 C CYS 136 61.688 80.557 1.00.337 1.00 53.47 ATOM 50 C CYS 136 61.688 80.557										1.00	
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ATOM 49 N CYS 136 58,954 79,191 -9,740 1.00 0.00 ATOM 50 H CYS 136 58,954 79,191 -9,740 1.00 0.00 55 ATOM 51 CA CYS 136 61.014 79,522 -9,495 1.00 57,62 ATOM 52 C CYS 136 61.688 80,552 -10,387 1.00 60,29 ATOM 53 O CYS 136 61,471 80,621 -11,594 1.00 62,12 ATOM 54 CB CYS 136 61,208 78,144 -10,115 1.00 53,99											
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ATOM 52 C CYS 136 61.688 80.552 -10.387 1.00 60.29 ATOM 53 O CYS 136 61.471 80.621 -11.594 1.00 62.12 ATOM 54 CB CYS 136 61.208 78.144 -10.115 1.00 53.99											
ATOM 53 O CYS 136 61.471 80.621 -11.594 1.00 62.12 ATOM 54 CB CYS 136 61.208 78.144 -10.115 1.00 53.99	55										60.29
ATOM 34 CB CYS 136 61.208 78.144 -10.115 1.00 53.99											
A10M 34 CB C13 58.70										1.00	
										1.00	58.70

	A 7503 /									
	ATOM	56	N	SER	137	62.538	81.402	-9.833	1.00	59.84
	MOTA	57	H	SER	137	62.572	81.519	-8.858	1.00	0.00
	ATOM	58	CA	SER	137	63.390	82.210	-10.667	1.00	62.00
	MOTA	59	СВ	SER	137	64.149	83.158			
5	ATOM	60	OG					-9.766	1.00	65.68
				SER	137	63.234	83.65 <i>5</i>	-8.792	1.00	75.16
	ATOM	61	HG	SER	137	62.492	84.111	-9.205	1.00	0.00
	MOTA	62	C	SER	137	64.313	81.329	-11.458	1.00	61,26
	ATOM	63	0	SER	137	64.602	80.202	-11.086	1.00	62.99
	ATOM	64	N	LEU	138	64.823				
10	ATOM	65	H	LEU			81.792	-12.585	1.00	61.53
	ATOM				138	64.686	82. 7 28	-12.829	1.00	0.00
		66	CA	LEU	138	65,553	80.911	-13.478	1.00	64 .28
	MOTA	67	CB	LEU	138	65.884	81.6 9 5	-14.748	1.00	63.19
	MOTA	68	CG	LEU	138	66.536	80.878	-15.866	1.00	61.50
	ATOM	69	CDI	LEU	138	65.823	79.540	-16.097	1.00	62.16
15	ATOM	70	CD2	LEU	138	66.528	81.749	-17.112	1.00	
	ATOM	71	c	LEU						65.07
	ATOM				138	66.813	80.309	-12.877	1.00	67.11
		72	0	LEU	138	67.183	79.164	-13.115	1.00	6 6.56
	ATOM	73	N	GLU	139	67.503	81.099	-12.063	1.00	68.85
	ATOM	74	H	GLU	139	67.248	82.038	-11.982	1.00	0.00
20	ATOM	75	CA	GLU	139	68.645	80.591	-11.330	1.00	71.48
•	ATOM	76	CB	GLU	139					
						69.271	81.757	-10.558	1.00	78.29
	MOTA	77	CG	GLU	139	68.277	82.677	-9.821	1.00	89.52
	ATOM	78	CD	GLU	139	68.983	83.966	-9.426	1.00	98.5
	ATOM	79	OE1	GLU	139	68.705	85.009	-10.033	1.00	99.9
25	ATOM	80	OE2	GLU	139	69.811	83.927	-8.510	1.00	
	ATOM	81	c	GLU						101.79
	ATOM				139	68.241	79.453	-10.411	1.00	69.73
		82	0	GLU	139	68.938	78.458	-10.328	1.00	70.27
	MOTA	83	N	GLU	140	67.107	79.556	-9.711	1.00	67.40
	MOTA	84	H	GLU	140	66.567	80.364	-9.814	1.00	0.00
30	ATOM	85	CA	GLU	140	66.616	78.489	-8.849	1.00	66.30
	MOTA	86	CB	GLU	140	65,290	78.874	-8.234	1.00	69.93
	MOTA	87	CG	GLU	140	65.411	80.248	-7.577	1.00	
	ATOM	88	CD	GLU						79.22
					140	64.097	80.745	-7.015	1.00	83.51
2 -	MOTA	89	OE!	GLU	140	63.207	79.947	-6.716	1.00	86.55
35	MOTA	90	OE2	GLU	140	63.971	81.956	-6 .866	1.00	89.13
	ATOM	91	С	GLU	140	66.431	77.221	-9.621	1.00	64.12
	ATOM	92	0	GLU	140	66.927	76.166	-9.273	1.00	61.28
	ATOM	93	N	ALA	141					
	ATOM	94	H			65.703	77.298	-10.720	1.00	64.66
4.0				ALA	141	65.236	78.135	-10.921	1.00	0.00
40	ATOM	95	CA	ALA	141	65.611	76.153	-11.604	1.00	68.98
	ATOM	96	CB	ALA	141	64.889	76.570	-12.884	1.00	70.25
	ATOM	97	С	ALA	141	66. 9 79	75.596	-11.947	1.00	71.14
	MOTA	98	0	ALA	141	67.313	74.428	-11.765	1.00	72.56
	ATOM	99	N	GLN	142	67.818			1.00	
45	ATOM	100	OH				76.487	-12.459		72.92
45				GLN	142	67.532	77.424	-12.537	1.00	0.00
	ATOM	101	CA	GLN	142	69 .151	76.115	-12.892	1.00	73.96
	ATOM	102	CB	GLN	142	69 .866	77.409	-13.316	1.00	77.97
	MOTA	103	CG	GLN	142	70.887	77.279	-14.452	1.00	87.44
	MOTA	104	CD	GLN	142	70.264	76.716	-15.714	1.00	92.53
50	ATOM	105	OEI	GLN	142	70.722	75.733	-16.286	1.00	95.95
-	MOTA	106								
			NE2	GLN	142	69.200	77.287	-16.242	1.00	93.80
	MOTA	107	HEZI	GLN	142	68.816	78.075	-15.810	1.00	0.00
	MOTA	108	HE22	GLN	142	68.852	76.862	-17.056	1.00	0.00
	MOTA	109	C	GLN	142	69.900	75.373	-11.802	1.00	71.44
55	ATOM	110	O	GLN	142	70.472		-12.010	1.00	
	ATOM	111					74.312			69.41
			N	ARG	143	69.911	75.911	-10.590	1.00	70.71
	MOTA	112	н	ARG	143	69.467	76.774	-10.440	1.00	0.00
	ATOM	113	CA	ARG	143	70.560	75.235	-9.482	1.00	71.58
	ATOM	114	CB	ARG	143	70.398	76.011	-8.169	1.00	66.09
60	ATOM	115	CG	ARG	143	71.452	77.103	-8.009	1.00	68.24
	ATOM	116	CD	ARG						
	ATOM	•			143	71.260	77.893	-6.715	1.00	67.96
	VIOW	117	NE	ARG	143	70.068	78.720	-6.772	1.00	68.10
	MOTA	118	RE	ARG	143	69.189	78.300	-6.871	1.00	0.00
	MOTA	119	CZ	ARG	143	70.158	80.048	-6.694	1.00	68.46

	ATOM	120	NH1	ARG	143	69.028	80.800	-6.754	1.00	68.32
	ATOM	121	HHII	ARG	143	68.137	80.358	-6.858	1.00	0.00 0.00
	ATOM	122	HH12	ARG	143	69.090	81.797	-6.697	1.00 1.00	68.45
	ATOM	123	NH2	ARG	143	71.365	80.664	-6.550 -6.498	1.00	0.00
5	MOTA	124	HH21	ARG	143	72.206	80.125	-6.490	1.00	0.00
	MOTA	125	HH22 .	ARG	143	71.412	81.661 73.850	-9.263	1.00	73.97
	MOTA	126	C	ARG	143 143	70.013 70.765	72.907	-9.074	1.00	77.08
	MOTA	127	O N	ARG ILE	144	68.696	73.662	-9.285	1.00	73.91
	MOTA	128	Н	ILE	144	68.105	74.411	-9.519	1.00	0.00
10	MOTA	129 130	CA	ILE	T44	68.143	72.352	-8.995	1.00	71.75
	ATOM	131	CB	ILE	144	66.605	72.394	-8.986	1.00	69.05
	ATOM	132	CG2	D.E	144	66.077	70.990	-8.729	1,00	69.68 65.22
	ATOM	133	CG1	ILE.	144	66.084	73.314	-7.895	1.00 1.00	62.44
15	ATOM	134	CDI	ILE	144	64.588	73.540	-8.061 10.036	1.00	72.66
	ATOM	135	C	ILE	144	68.604	71.367	-10.036 -9.743	1.00	71.17
	MOTA	136	٥	ILE.	144	69.045	70.260 71.765	-11.304	1.00	75.70
	MOTA	137	N	TRP	145	68.507 68.167	72.662	-11.514	1.00	0.00
	MOTA	138	H	TRP	145 145	68.929	70.861	-12.351	1.00	83.94
20	MOTA	139	CA CB	TRP TRP	145	68.658	71.462	-13.746	1.00	86.15
	MOTA MOTA	140 141	CG	TRP	145	69.038	70.460	-14.839	1.00	93.70
	MOTA	142	CD2	TRP	145	68.217	69,255	-15.098	1.00	95.01
	ATOM	143	CE2	TRP	145	69.056	68.711	-16.236	1.00	96.30
25	ATOM	144	CE3	TRP	145	67.071	68.594	-14.711	1.00	94.15 95.29
	ATOM	145	CD1	TRP	145	70.133	70.564	-15.670	1.00 1.00	93.47
	ATOM	146	NEI	TRP	145	70.111	69,520	-16.475 -17.166	1.00	0.00
	ATOM	147	HEI	TRP	145	70.787	69.359 67.562	-16.873	1.00	100.90
	ATOM	148	CZ2	TRP	145	68.642 66.717	67.444	-15.394	1.00	96.37
30	MOTA		CZ3	TRP TRP	145 145	67,473	66,945	-16.445	1.00	98.93
	ATOM		CH2	TRP	145	70.411	70.588	-12.184	1.00	87.50
	MOTA MOTA		C O	TRP	145	70.817	69.442	-12.042	1.00	88.19
	ATOM		N	ALA	146	71.240	71.638	-12.196	1.00	90.23
35	ATOM		H	ALA	146	70.872	72.546	-12.279	1.00	0.00 90.96
55	ATOM		CA	ALA	146	72.678	71.479	-12.055	1.00 1.00	89.47
	ATOM		CB	ALA	146	73.316	72.866	-12.003	1.00	92.53
	ATOM		С	ALA.	146	73.078	70.678	-10.827 -10.833	1.00	93.84
	ATOM		0	ALA	146	74.075	69.975 70.767	-9.739	1.00	93.24
40				GLN	147	72.310 71.578	71.421	-9.711	1.00	0.00
	MOTA			GLN	147 147	72.543	69.925	-8.583	1.00	94.09
	YLO?		_	GLN GLN	147	71.690	70.379	-7.387	1.00	96.74
	ATON			GLN	147	72.033	69.642	-6.090	1.00	102.36
45	AOTA AOTA			GLN	147	70.829	69.585	-5.181	1.00	107.75
	KOTA			GLN	147	69.729	70.018	-5.491	1.00	109.80
	ATO			GLN	147	70.957	68.991	-4.006	1.00	109.34 0.00
	ATO!			GLN	147	71.821	68.602	-3.760	1,00 1,00	0.00
	ATO				147	70.155	68.987	-3.438 9.875	1.00	93.64
50			C	GLN	. 147	72.216	68.479 67.595	-8.875 -8.801	1.00	93.43
	ATO			GLN	147	73.058 70. 96 9	68.171	-9.211	1.00	91.82
	ATO			LYS	148	70.319	68.869	-9.439	1.00	0.00
	ATO			LYS	148 148	70.621	66.770	-9.239	1.00	89.79
_	ATO			LYS LYS	148	69.339	66.539	-8.438	1.00	91.49
5				LYS	148	69.449	66.861	-6.954	1.00	91.59
	ATO			LYS	148	68.211	66.386	-6.193	1.00	94.76
	ATO			LYS	148	68.288	66.715	-4.706	1.00	97,17
	ATO ATO			LYS	148	69.521	66.190	4.145	1.00	102.16
6				LYS	148	70.339	66.615	-4.627	1.00	0.00 0.00
•	ATO		0 HZ2		148	69.553		-4.275	1.00	0.00
	ATO		81 HZ3		148			-3.131	1.00 1.00	87.23
	ATO		82 C	LYS	148			-10.578 -10.625	1.00	88.22
	ATO		83 O	LYS	148	70.294	64,882	-10.023	1.00	

	MOTA	184	N	ALA	149	70.496	66.825	-11.691	1.00	83.64
	MOTA	185	H	ALA	149	70.728	67.775	-11.599	1.00	0.00
	ATOM	186	CA	ALA	149	70.271	66.316	-13.039	1.00	81.28
_	ATOM	187	СВ	ALA	149	71.416	66.848	-13.906	1.00	80.46
5	ATOM	188	С	ALA	149	70,092	64.807	-13.272	1.00	80.23
	ATOM	189	0	ALA	149	69.034	64.377	-13.709	1.00	81.20
	ATOM	190	N	ALA	150	71.079	63.9 39	-13.002	1.00	77.26
	MOTA	191	H	ALA	150	71.903	64,290	-12.604	1.00	0.00
7.0	ATOM	192	CA	ALA	150	70.928	62.493	-13.224	1.00	72.54
10	ATOM	193	СВ	ALA	150	72.254	61.803	-12.872	1.00	71.93
	ATOM	194	0 . C	ALA	150	69.786	61.803	-12.458	1.00	67.17
	MOTA MOTA	195 196		ALA	150	69.332	60.728	-12.832	1.00	64.24
	ATOM	197	N H	GLU	151	69.298	62.391	-11.366	1.00	58.90
15	ATOM	198	ČΑ	GLU GLU	151	69.693	63.225	-11.031	1.00	0.00
	ATOM	199	CB	GLU	151	68.153	61.823	-10.700	1.00	55.46
	ATOM	200	CG	GLU	15t 15t	68.612	61.333	-9.316 0.533	1.00	60.03
	ATOM	201	CD	GLU	151	69.510 69.733	60.091	-9.532	1.00	67.91
	ATOM	202	OE1	GLŲ	151	68.753	59.292 58.778	-8. 2 59 -7.702	1.00 1.00	72.08
20	ATOM	203	OE2	GLU	151	70.887	59.159	-7.702 -7.846		75.94
	ATOM	204	c	GLU	151	66.966	62.759	-10.610	1.00	70.44
	ATOM	205	ō	GLU	151	66.237	62.811	-9.632	1.00	49.03
	ATOM	206	N	ILE	152	66.751	63.538	-9.632 -11.671	1.00 1.00	40.75 46.24
	ATOM	207	H	ILE	152	67.390	63.525	-12.419	1.00	0.00
25	ATOM	208	CA	ILE	152	65.578	64.379	-11.820	1.00	41.23
	MOTA	209	CB	ILE	152	66.006	65.876	-11.979	1.00	41.97
	MOTA	210	CG2	ILE	152	64.846	66.776	-12.375	1.00	37.44
	MOTA	211	CGI	ILE	152	66.456	66.419	-10.633	1.00	39.50
20	MOTA	212	CDI	IL.E	152	67.026	67.826	-10.790	1.00	45.38
30	MOTA	213	C ·	ILE :	152	64.838	63.880	-13.063	1.00	39.41
	MOTA	214	0	ILE	152	65.399	63.521	-14.089	1.00	38.70
	MOTA MOTA	215	N	TYR	153	63.506	63.846	-12.966	1.00	37.18
	ATOM	216 217	H	TYR	153	63.125	64.015	-12.080	1.00	0.00
35	ATOM	218	CA	TYR	153	62.606	63.601	-14.098	1.00	34.02
22	ATOM	219	CB CG	TYR TYR	153	61.131	63.521	-13.608	1.00	30.99
	ATOM	220	CD1	TYR	153 153	60.815	62.157	-13.006	1.00	28.21
	ATOM	221	CE1	TYR	153	60.889 60.661	61.023 59.769	-13.795 -13.266	00.1 00.1	27.24
	ATOM	222	CD2	TYR	153	60.496	62.024	-11.663	1.00	25.65 24.92
40	ATOM	223	CE2	TYR	153	60.266	60.774	-11.126	1.00	20.91
	ATOM	224	CZ	TYR	153	60.352	59.657	-11.934	1.00	24.92
	ATOM	225	OH	TYR	153	60.116	58.405	-11.404	1.00	22.74
	MOTA	226	HH	TYR	1.53	59.896	58.542	-10.481	1.00	0.00
	ATOM	227	С	TYR	153	62.734	64.754	-15.092	1.00	31.41
45	ATOM	228	0	TYR	153	62.545	65.895	-14.708	1.00	30.52
	ATOM	229	N	PRO	154	63.045	64.585	-16.332	1.00	29.09
	ATOM	230	CD	PRO	154	63.433	63.298	-16.886	1.00	26.14
	ATOM	231	CA	PRO	154	63.119	65.685	-17.300	1.00	26.59
50	MOTA MOTA	232 233	CB	PRO	154	63.791	65.078	-18.508	1.00	21.73
50	ATOM	233 234	CG	PRO	154	63,408	63.614	-18.383	1.00	26.66
	MOTA	235	С 0	PRO	154	61.782	66.309	-17.627	1.00	27.17
	ATOM	236	N	PRO ILE	154	60.763	65.693	-17.860	1.00	30.28
	ATOM	237	H	ILE	155 155	61.791 62.636	67.620	-17.645	1.00	27.49
55	ATOM	238	CA	n.e	155	60.623	68.068 68.417	-17.453	1.00	0.00
	ATOM	239	CB	ILE	155	60.810		-17.912	1.00	27.05
	ATOM	240	CG2	ILE	155	59.893	69.687 70.851	-17.067 -17.429	1.00 1.00	27.13 28.62
	ATOM	241	CGI	ILE	155	60.584	69.235	-17.429	1.00	28.62 27.98
	ATOM	242	CDI	ILE	155	59.370	68.340	-15.354	1.00	30.26
60	MOTA	243	C	ILE	155	60.648	68.636	-19.414	1.00	33.41
	ATOM	244	0	ILE	155	61.715	68.804	-19.990	1.00	34.72
	ATOM	245	N	MET	156	59.510	68.638	-20.116	1.00	33.95
	MOTA	245	н	MET	156	58.677	68.455	-19.642	1.00	0.00
	MOTA	247	CA	MET	156	59,471	68.939	-21.542	1.00	31.70

											21.01
	ATOM	248	CB	ME	ΕT	156	58.252	68.339	-22.224	1.00 1.00	31.01 30.35
	ATOM	249	CG	ME		156	58.574	67.001	-22.838 -23.577	1.00	39.82
	MOTA	250	SD	ME		156	57.118	66.252 64.545	-23.528	1.00	34.89
	MOTA	251	CE	M		156	57.590 59.384	70.434	-21.735	1.00	34.78
5	MOTA	252	C	MI		156	59.035	71.208	-20.854	1.00	32.33
	MOTA	253	0	MI		156 157	59.711	70.869	-22.942	1.00	37.95
	MOTA	254	N		SP SP	157	59.923	70.224	-23.648	1.00	0.00
	MOTA	255 256	H CA		SP	157	59.715	72.281	-23.240	1.00	38.36
10	MOTA MOTA	257	CB		SP	157	60.385	72.438	-24.613	1.00	51.68
10	MOTA	258	CG		SP.	157	60.369	73.848	-25.178	1.00	61.42 68.30
	ATOM	259	ODI		.SP	157	60.483	73.957	-26.401	1.00 1.00	67.14
	ATOM	260	OD2	A	SP	157	60.249	74.822	-24.421	1.00	31.15
	MOTA	261	C		.SP	157	58.315	72.829	-23.203 -23.730	1.00	27.50
15	MOTA	262	0		SP	157	57,360	72.284 73. 9 65	-22.543	1.00	30.63
	MOTA	263	N		.YS	158	58.232 59.032	74.318	-22.102	1.00	0.00
	MOTA	264	H		.YS	158 158	56.975	74.638	-22.320	1.00	33.69
	ATOM	265	CA		.YS .YS	158	57.265	75.906	-21.540	1.00	34.01
	ATOM	266	CB CG		.13 .YS	158	56.037	76.664	-21.085	1.00	40.80
20	ATOM ATOM	267 268	CD		LYS	158	56.390	77.826	-20.154	1.00	48.58
	ATOM	269	CE		LYS	158	55.152	78.472	-19.524	1.00	52.61
	ATOM	270	NZ		LYS	158	55.537	79.342	-18.431	1.00	59.47 0.00
	ATOM	271	HZ	1 1	LYS	158	56.154	80.095	-18.794	1.00 1.00	0.00
25	ATOM	272	HZ		LYS	158	56.050	78.792	-17.712 -17.998	1.00	0.00
	ATOM	273	HZ:		LYS	158	54.691	79.763 74.942	-23.618	1.00	38.44
	ATOM		c		LYS	158	56.275 55.076	74.803	-23.735	1.00	41.75
	MOTA		0		LYS	158 159	56.962	75.367	-24.672	1.00	41.20
	MOTA				SER SER	159	57.938	75.470	-24.645	1.00	0.00
30	MOTA MOTA		H CA		SER	159	56.297	75.717	-25.916	1.00	39.99
	ATOM				SER	159	57.304	76.202	-26.930	1.00	39.69 40.47
	ATOM				SER	159	58.553	75.552	-26.729	1.00 1.00	0.00
	ATOM				SER	159	58.453	74.606	-26.880 36.548	1.00	40.83
35	ATOM		C		SER	159	55.543	74.590	-26.548 -27.370	1.00	43.09
	MOTA	283			SER	159	54.666	74.774 73.373	-26.184	1.00	43.73
	ATON				SER	160	55.886 56.444	73.181	-25.399	1.00	0.00
	ATOM				ser ser	160 160	55.410	72.261	-26.959	1.00	45.71
4.5	ATON				SER	160	56.592	71.756	-27.769	1.00	45.15
40					SER	160	57.835	71.958	-27.083	1.00	53.30
	ATON MOTA				SER	160	58.517	71.533	-27.614	1.00	0.00 45.78
	ATO	-		~	SER	160	54.800	71.154	-26.121	1.00	48.03
	ATO	-			SER	160	54.262	70.217	-26.700	1.00 1.00	38.45
45					ARG	161	54.842	71.162	-24.782 -24.278	1.00	0.00
	ATO				ARG	161	55.201	71.921 70.004	-24.097	1.00	32.89
	ATO	M 29		A.	ARG	161	54.322	69.929	-22.727	1.00	31.71
	ATO		_	В	ARG	161	54.987 54.631	71.054	-21.769	1.00	29.23
	OTA			G	ARG	161 . • 161	55.579	71.078	-20.571	1.00	26.52
50				:D	ARG	161	55.148	72.182	-19.760	1.00	23.86
	ATO			(E	ARG ARG	161	54.189	72.357	-19.663	1.00	0.00
	ATO			E Z	ARG	161	56.001	72.971	-19.138	1.00	24.88
	ATO			H1	ARG	161	55.486		-18.477	1.00	23.22 0.00
5				ihii	ARG	161	54,499		-18.481	1.00	0.00
9	ATO	-		HH12	ARG	161	56.093		-18.006	1.00	26.60
	ATO			NHZ	ARG	161	57.339		-19.115	1.00	0.00
	TA			HH21	ARG	161			-19.572		0.00
	ATO			HH22	ARG	161					30.40
6	O AT			С	ARG	161				1.00	30.63
·	AT		308	O	ARG	161					25.20
	AT			N	THR	162					
	AT			H	THR	162					
	AT	OM :	311	CA	THR	163	٧٥٠.٥٧	, 00.70.			

- 47 -

	ATOM	312	CB	THR	162	50.068	40 226	06.000		
	ATOM	313	OGI	THR	162	50.780	68.276	-25.028	1.00	23.65
	ATOM	314	HG!	THR	162	50.721	67.056	-25.252	1.00	26.56
	MOTA	315	CG2	THR	162		66.510	-24.445	1.00	0.00
5	ATOM	316	c	THR	162	50.148	69.114	-26.306	1.00	23.19
_	MOTA	317	ō	THR		50.373	68.219	-22.510	1.00	25.37
	ATOM	318	N	ARG	162	50.062	67.030	-22.548	1.00	27.75
	ATOM	319	н		163	50.423	68.826	-21.340	1.00	24.62
	ATOM	320		ARG	163	50.560	69.79 7	-21.312	1.00	0.00
10	MOTA	321	CA	ARG	163	50.174	68.133	-20.077	1.00	24.12
20	MOTA	322	CB CG	ARG	163	50.757	68.905	-18.918	1.00	15.48
	ATOM	323		ARG	163	52.227	68.612	-18.793	1.00	17.34
	ATOM		CD	ARG	163	52.721	69.593	-17.772	1.00	18.83
		324	NE	ARG	163	54.136	69.427	-17.669	1.00	24.11
15	MOTA	325	HE	ARG	163	54.592	68.736	-18.195	1.00	0.00
13	MOTA	326	CZ	ARG	163	54.822	70.232	-16.883	1.00	25.67
	ATOM	327	NHI	ARG	163	56.160	70.051	-16.807	1.00	34.46
	ATOM	328	HHII	ARG	163	56.596	69.318	-17.328	1.00	0.00
	ATOM	329	HH12	ARG	163	56.710	70.634	-16.209	1.00	0.00
	MOTA	330	NH2	ARG	163	54.219	71.220	-16.167	1.00	27.39
2,0	MOTA	331	HH21	ARG	163	53.229	71.357	-16.223	1.00	0.00
	MOTA	332	HH22	ARG	163	54.772	71.806	-15.575	1.00	0.00
	MOTA	333	С	ARG	163	48.738	67.876	-13.373	1.00	
	MOTA	334	Ō	ARG	163	47.893	68.762	-19.714		24.62
	MOTA	335	N	LEU	164	48.357			1.00	29.04
25	MOTA	336	н	LEU	164	49.023	66.667 65.971	-19.340	1.00	23.75
	ATOM	337	CA	LEU	164	46.944		-19.147	1.00	0.00
	ATOM	338	CB	LEU	164	46.366	66.455	-19.043	1.00	24.57
	MOTA	339	CG	LEU	164		65.336	-19.893	1.00	27.72
	ATOM	340	CDI	LEU	164	45.556	65.607	-21.165	1.00	28.04
30	ATOM	341	CD2	LEU	164	46.108	66.789	-21.926	1.00	30.94
	ATOM	342	C	LEU	164	45.573	64.331	-22.006	1.00	27.16
	ATOM	343	ŏ	LEU	164	46.774	66.047	-17.612	1.00	23.76
	ATOM	344	N	ALA	165	47.607	65.281	-17.142	1.00	22.71
	ATOM	345	H	ALA		45.733	66.505	-16.907	1.00	21.51
35	MOTA	346	CA	ALA	165	45.124	67.175	-17.292	1.00	0.00
	ATOM	347	CB	ALA	165	45.396	65.986	-15.589	1.00	24.14
	ATOM	348	C	ALA	165	45.687	67.006	-14.488	1.00	21.16
	ATOM	349	ō	ALA	165	43.924	65.668	-15.543	1.00	24.40
	ATOM	350	N		165	43.162	66.279	-16.279	1.00	25.81
40	ATOM	351	H	LEU	166	43.495	64.730	-14.699	1.00	24.86
30	ATOM	352	CA	LEU	166	44.143	64.282	-14.114	1.00	0.00
	ATOM	352 353		LEU	166	42.091	64.361	-14.592	1.00	27.92
			CB	LEU	166	41.914	62.948	-15.196	1.00	23.03
	MOTA	354	CG	LEU	166	40.571	62.280	-14.919	1.00	20.45
45	ATOM	355	CDI	LEU	166	39.446	62.990	-15.658	1.00	14.43
43	ATOM	356	CD2	LEU	166	40.691	60.807	-15.320	1.00	21.71
	ATOM	357	C	LEU	166	41.580	64.404	-13.149	1.00	28.64
	ATOM	358	0	LEU	166	42.207	63.838	-12.261	1.00	26.66
	MOTA	359	N	ILE	167	40.441	65.079	-12.901	1.00	27.07
	MOTA	360	H	ILE	167	39.991	65.549	-13.638	1.00	0.00
50	ATOM	361	CA	ILE	167	39.802	65.137	-11.600	1.00	21.18
	ATOM	362	CB	ILE	167	39.494	66.602	-11.205	1.00	20.90
	ATOM	363	CG2	ILE	167	38.719	66.663	-9.893	1.00	21.77
	ATOM	364	CG1	ILE	167	40.793	67.354	-11.003	1.00	17.47
	ATOM	365	CD1	ILE	167	40.499	68.753	-10.522	1.00	17.98
55	MOTA	366	C	TLE.	167	38.513	64.358	-11.684	1.00	24.82
	ATOM	367	0	ILE	167	37.634	64.653	-12.493	1.00	23.12
	ATOM	368	N	ILE	168	38.333	63.328	-10.865	1.00	25.26
	ATOM	369	H	ILE	168	39.066	63.004	-10.297	1.00	0.00
	ATOM	370	CA	ILE	168	37.022	62.731			
60	ATOM	371	СВ	ILE	168	37.119	61.228	-10.757 -11.076	1.00 1.00	23.44 23.97
-	ATOM	372	CG2	ILE	168	35.741	60.582			
	ATOM	373	CG1	ILE	168	37.581	61.030	-10.863	1.00	28.80
	ATOM	374	CDI	ILE	168			-12.546	1.00	22.39
	ATOM	375	C	ILE		37.869	59.587	-12.959	1.00	22.21
	1110011	5,5	<u> </u>	تلبط	168	36.506	62.981	-9.353	1.00	24,34

										23.09
	ATOM	376	0	ILE	168	37.126	62.633	-8.355 -9.260	1.00 1.00	26.60
	ATOM	377	N	CYS	169	35.337	63.613 63.777	-10.079	1.00	0.00
	ATOM	378	H	CYS	169 169	34.821 34.765	64.030	-7.986	1.00	27.82
_	ATOM	379	CA	CYS CYS	169	34.847	65.544	-7.831	1.00	26.82
5	ATOM	380 381	CB SG	CYS	169	34.147	66.177	-6.282	1.00	31.50
	ATOM ATOM	382	C	CYS	169	33.323	63.629	-7.866	1.00	28.36
	ATOM	383	ō	CYS	169	32.520	63.950	-8.735	1.00 1.00	25.83 27.23
	ATOM	384	N	ASN	170	32,948	62.918	-6.803 -6.124	1.00	0.00
10	ATOM	385	H	ASN	170	33.617	62.683 62.555	-6.619	1.00	29.76
	ATOM	386	CA.	NZA	170	31.554 31.372	61.071	-6.286	1.00	26.22
	ATOM	387	CB.	ASN ASN	170 170	31.674	60.138	-7.440	1.00	29.16
	ATOM	388	CG OD1	ASN	170	31.709	60.485	-8.614	1.00	32.47
15	ATOM ATOM	389 390	ND2	ASN	170	31.970	58.887	-7.151	1.00	25.45
13	ATOM	391	HD21	ASN	170	32.015	58.569	-6.216	1.00	0.00 0.00
	ATOM	392	HD22	ASN	170	32.132	58.288	-7.901	1.00	32.01
	ATOM	393	C	ASN	170	30.957	63.328	-5.478 -4.361	1.00	34.71
	ATOM	394	0	ASN	170	31.382	63.086 64.247	-5.672	1.00	36.94
20	ATOM	395	N	GLU	171 171	30.000 29.711	64.420	-6.594	1.00	0.00
	ATOM	396	H	GLU GLU	171	29.301	64.905	-4.558	1.00	39.76
	ATOM	397 398	CA CB	GLU	171	29.119	66.413	-4.822	1.00	36.97
	MOTA MOTA	399	CG	GLU	171	28.418	67.084	-3.634	1.00	43,98 47,95
25	ATOM	400	CD	GLU	171	28.233	68.579	-3.777	1.00 1.00	53.49
	ATOM	401	OE1	GLU	171	28.302	69.095	-4.890 -2.761	1.00	48.17
	ATOM	402	OE2	GLU	171	28.003	69.233 64.338	-4.240	1.00	41.90
	MOTA	403	C	GLU	171 171	27.914 27.509	64.214	-3.096	1.00	40.97
	ATOM	404	0	GLU GLU	172	27.133	63.976	-5.251	1.00	48.24
30	ATOM	405 406	N H	GLU	172	27.505	63.929	-6.159	1.00	0.00
	MOTA MOTA		ČA	GLU	172	25.760	63.534	-5.050	1.00	51.31
	MOTA		CB	GLU	172	24.802	64.170	-6.061	1.00 1.00	55.01 66.24
	ATOM		CG	GLU	172	23.819	65.134	-5.408 -5.699	1.00	74.35
35	ATOM	410	CD	GLU	172	24.170	66.580 67.402	-5.708	1.00	81.11
	ATOM		OE1	GLU	172	23.249 25.342	66.893	-5.911	1.00	76.04
	ATOM		OEZ	GLU GLU	172 172	25.676	62.035	-5.242	1.00	49.78
	MOTA MOTA		С 0	GLU	172	26.092	61.501	-6.266	1.00	46.42
40	ATOM		N	PHE	173	25.125	61.338	-4.250	1.00	49,39 0.00
-30	ATOM		Н	PHE	173	24. 69 5	61.780	-3.485	1.00 1.00	53.47
	ATOM		CA	PHE	173	25.085	59.888	-4.303 -3.142	1.00	50.77
	ATOM	418	CB	PHE	173	25.878	59.313 59.547	-3.286	1.00	46.81
_	ATON			PHE	173	27.371 28.173	58.521	-3.768	1.00	40.36
45				PHE PHE	173 173	27.927	60.747	-2.867	1.00	42.17
	ATON			PHE	173	29.535	58.684	-3.778	1.00	37,00
	IOTA IOTA			PHE	173	29.291	60.904	-2.902	1.00	38.90 39.65
	OTA			PHE	173	30,089	59.870	-3.332	1.00	56.41
50				PHE	. 173	23,661	59.372	-4.221 -3.675	1.00 1.00	60.38
•	ATO		5 0	PHE	173	22,772	60.010 58.194	-3.673 -4.764	1.00	54.59
	ATO			ASP	174	23.394 24.131	57.657	-5.134	1.00	0.00
	ATO			ASP ASP	174 174	22.044	57.658	4.745	1.00	53.81
	ATO			ASP	174	21.943	56.370	-5.557	1.00	51.10
5				ASP	174	22.091	56.666	·7.030	1.00	53.87
	OTA OTA				174	22.150	55.698	-7.795	1.00	53.10 53.80
	OTA				174	22.148		-7.408 -7.408	1.00 1.00	56.79
	OTA		-	ASP	174	21.500		-3,370 -3.086	1.00	60.64
6			15 O	ASP	174	20.317		-3.060 -2.457	1.00	57.04
	ATC			SER	175				1.00	0.00
	ATO			SER	175 175				1.00	56.82
	ATO		38 CA	SER	175				1.00	53.73
	λTO	JM 4	39 CB	SER	.,,					

	MOTA	440	OG	SER	175	21.821	54.307	-2.423	1.00	50.93
	ATOM	441	HG	SER	175	22.453	54.685	-3.041	1.00	0.00
	MOTA	442	c	SER	175	22.614	57.115	-0.038	1.00	59.19
	MOTA	443	0	SER	175	22.050	57.584	0.945	1.00	63.80
5	MOTA	444	N	ILE	176	23.949	57.172	-0.133	1.00	57.13
	ATOM	445	H	ILE	176	24.384	56.889	-0.963	1.00	0.00
	ATOM	446	CA	ILE	176	24.725	57.758	0.942	1.00	50.84
	MOTA	447	CB	ILE	176	26.167	57.155	0.920	1.00	52.16
	ATOM	448	CG2	ILE	176	25.982	55.645	1.127	1.00	54.09
10	MOTA	449	CGI	ILE	176	26.947	57.381	-0.375	1.00	51.82
	MOTA	450	CDI	ILE	176	28.205	56.486	-0.451	1.00	47.41
	ATOM	451	c ·	ILE	176	24.716	59.262	0.770	1.00	47.11
	ATOM	452	ŏ	ILE	176	24,516	59.755	-0.332	1.00	45.21
	ATOM	453	N	PRO	177	24.918	60.009	1.801	1.00	47.09
15	ATOM	454	CD	PRO	177	25.067	59.502	3.162	1.00	46.36
	ATOM	455	CA	PRO	177	24.821	61.475	1.806	1.00	48.52
	MOTA	456	CB	PRO	177	24.733	61.862	3.280	1.00	45.72
	ATOM	457	CG	PRO	177	25.513	60.744	3.937	1.00	45.89
	ATOM	458	Č	PRO	177	25.870	62.308	1.096	1.00	
20	ATOM	459	ō	PRO	177	27.078		1.042		51.58
-:	ATOM	460	Ň	ARG	178		62.045		1.00	56.38
	ATOM	461	н	ARG	178	25.287	63.373	0.559	1.00	50.95
	ATOM	462	CA	ARG		24.351	63.552	0.778	1.00	0.00
	ATOM	463			178	25.959	64.288	-0.328	1.00	52.24
25	ATOM	464	CB CG	ARG	178	24.910	65.270	-0.842	1.00	54.80
23	ATOM	465	CD	ARG	178	25.505	66.396	-1.661	1.00	62.55
	ATOM	466	NE	ARG ARG	178	24.484	67.300	-2.273	1.00	67.32
	ATOM	467	HE	ARG	178	25.167 26.144	68.507	-2.664	1.00	77.79
	MOTA	468	cz	ARG	178 178		68.522 69.607	-2.727	1.00	0.00
30	ATOM	469	NH1	ARG	178	24.473		-2.918	1.00	85.85
	ATOM	470	HH11	ARG	178	23.102 22.609	69.585 68.736	-2.899 -2.708	1.00 1.00	92.67 0.00
	ATOM	471	HH12	ARG	178	22.586	70.415	-3.103		0.00
	MOTA	472	NH2	ARG	178	25.133	70.413	-3.165	1.00	90.17
	ATOM	473	HH21	ARG	178	26.133	70.802		1.00	
35	ATOM	474	HH22	ARG	178	24.609		-3.145	1.00	0.00
	ATOM	475	C .	ARG	178		71.605	-3.354	1.00	0.00
	MOTA	476	ŏ	ARG	178	27.133	65.020	0.280	1.00	49.82
	ATOM	477	N	ARG	179	27.078	65.589	1.354	1.00	51.03
	ATOM	478	н	ARG	179	28.243 28.218	65.007	-0.439	1.00	47.03
40	ATOM	479	CA	ARG	179		64.607	-1.336	1.00	0.00
. •	ATOM	480	CB CB	ARG	179	29.458	65.625	0.039	1.00	42.75
	ATOM	481	CG	ARG	179	30.625 30.599	64.919 63.395	-0.632	1.00	42.27
	ATOM	482	CD	ARG	179	31.759	62.682	-0.447 -1.159	1.00	39.14 40.61
	ATOM	483	NE	ARG	179	33.057	63.164	-0.704	1.00	40.03
45	ATOM	484	HE	ARG	179	33.387	62.869	0.169	1.00	0.00
	ATOM	485	CZ	ARG	179	33.827°	63.996	-1.427	1.00	40.60
	MOTA	486	NHI	ARG	179	33.483	64,391	-2.691	1.00	35.23
	ATOM	487	HHII	ARG	179	32.630	64.073	-3.103	1.00	0.00
	ATOM	488	HH12	ARG	179	34,077	65.023	-3.191	1.00	0.00
50	ATOM	489	NH2	ARG	179	34.965	64.483	-0.859	1.00	35.40
	ATOM	490	HHZI	ARG	179	35.205	64.214	0.074	1.00	0.00
	ATOM	491	HH22	ARG	179	35.562	65.100	-1.370	1.00	0.00
	ATOM	492	C	ARG	179	29.512	67.124	-0.198	1.00	44.55
	ATOM	493	Ō	ARG	179	30.215	67.124			
55	ATOM	494	N	THR	180	28.730	67.862	-1.034 0.570	1.00 1.00	44.42 45.86
	ATOM	495	н	THR	180	28.0G3				
	MOTA	496	CA.	THR			67.420	1.138	1.00	0.00
	MOTA	497	CB	THR	180	28.816	69.308	0.548	1.00	46.36
	MOTA	498	OGI	THR	180 180	27.770	69.840	1.532	1.00	50.60
60	ATOM	499	HG1	THR		26.517	69.456	0.968	1.00	54.63
- 0	ATOM	500	CG2	THR	180	25.800 27.792	69.689	1.568	1.00	0.00
	ATOM	501			180		71.351	1.735	1.00	53.13
	ATOM	502	С 0	THR THR	180	30.221	69.783	0.901	1.00	46.41 47.03
	ATOM	503	N	GLY	180 181	30.882	69.285	1.807	1.00	47.93 46.30
			**	ar 1	101	30.713	70.783	0.171	1.00	46.30

						00.000	71.312	-0.350	1.00	0.00
	ATOM	504	H	GLY	181	30.090 32.118	71.193	0.236	1.00	41.25
	MOTA	505	CV	GLY	181	32.915	70.679	-0.948	00.1	40.64
	MOTA	506	C	GLY	181	34.005	71.154	-1.278	1.00	37.29
_	MOTA	507	0	GLY	181 182	32.381	69.688	-1.672	1.00	35.93
5 -	MOTA	508	И	ALA	182	31.540	69.257	-1.418	1.00	0.00
	ATOM	509	H	ALA ALA	182	33.136	69.171	-2.785	1.00	39.67
	ATOM	510	CA	ALA	182	32.314	68.028	-3.377	1.00	35.68
	ATOM	511	CB C	ALA	182	33.503	70.220	-3.824	1.00	39.67
10	ATOM	512 513	ò	ALA	182	34.590	70.239	4.378	1.00	40.75
10	MOTA MOTA	514	N	GLU	183	32.599	71.153	-4.107	1.00	42.72
	ATOM	515	H	GLU	183	31.748	71.149	-3.627	1.00	0.00
	ATOM	516	Ċλ	GLU	183	32.824	72.180	-5.108	1.00	40.12
	ATOM	517	CB	GLU	183	31.614	73.091	-5.246	1.00	47.30 63.74
15	ATOM	518	CG	GLU	183	31.637	73.914	-6.544	1.00	71.48
	ATOM	519	CD	GLU	183	30.918	73.216	-7.697	1.00 1.00	78.62
	MOTA	520	OE1	GIL U	183	30.272	73.920	-8.481	1.00	76.00
	MOTA	521	OE2	GLU	183	30.994	71.988	-7.818 -4.776	1.00	35.44
	ATOM	522	С	GLU	183	33.998	73.045	-5.639	1.00	37.02
20	MOTA	523	0	GLU	183	34.778	73.390 73.424	-3.521	1.00	32.69
•	MOTA	524	N	VAL	184	34.197	73.236	-2.857	1.00	0.00
	MOTA	525	H	VAL	184	33.502 35.437	74.093	-3.114	1.00	32.56
	MOTA	526	CA	YAL	184 184	35,391	74.286	-1.598	1.00	30.19
	ATOM	527	CB	VAL VAL	184	36.568	75.090	-1.054	1.00	28.08
25	MOTA	528	CGI	VAL	184	34.081	74.992	-1.308	1.00	34.05
	ATOM	529 530	CG2 C	VAL	184	36.707	73.335	-3.505	1.00	31.95
	MOTA MOTA	531	ō	VAL	184	37.716	73.836	-3.987	1.00	31.94
	ATOM	532	N	ASP	185	36.649	72.025	-3.278	1.00	36.44
30	ATOM		H .	ASP	185	35.811	71.642	-2.945	1.00	0.00
50	ATOM		CA	ASP	185	37.768	71.151	-3.586	1.00	32.53 33.96
	ATOM		CB	ASP	185	37.470	69.728	-3.077	1.00	35.35
	ATOM		CG	ASP	185	37.375	69.592	-1.539	1.00	28.61
	ATOM		0D1	ASP	185	38.047	70.367	-0.833	1.00	32.20
35	MOTA		OD2	ASP	185	36.635	68.705	-1.082 -5.059	1.00	30.28
	ATOM	539	С	ASP	185	38.020	71.139 71.233	-5.493	1.00	29.51
	ATOM		0	ASP	185	39.161	71.032	-5.849	1.00	29.28
	ATOM		N	ILE	186	36.956 36.070	70.902	-5.447	1.00	6.00
	ATOM		H	ILE	186 186	37.096	71.078	-7.305	1.00	31.19
40			CA	ile Ile	186	35.728	70.861	-8.016	1.00	31.31
	ATOM		CB CG2	ILE	186	35.874	71.027	-9.544	1.00	29.98
	ATON		CGI	TLE	186	35.212	69.446	-7.688	1.00	33.52
	KOTA KOTA		CDI	ILE	186	33.829	69.127	-8.292	1.00	24.25
45			c .	ILE	186	37.684	72.413	-7.748	1.00	30.16
7.0	ATOL	-	ō	ILE.	186	38.691	72,466	-8.447	1.00	30.56 27.47
	ATO			THR	187	37.132	73.560	-7.388	1.00	0.00
	ATO			THR	187	36.310	73.583	-6.842	1.00 1.00	27.93
	ATO			THR	187	37.739	74.820	-7.788 7.176	1.00	27.45
50) ATO			THR	. 187	36.943	75.985	-7.175 -7.625	1.00	32.61
	ATO:	M 554	OGI	THR	187	35.600	75.803	-7.023 -8.583	1.00	0.00
	ATO	M 555	HG1	THR	187	35.578	75.839	-7.566	1.00	24.19
	ATO	M 556	CG2	THR	187	37.461	77.371 74.935	-7.300 -7.395	1,00	28.16
	ATO	M 557		THR	187	39.193	75.448	-8.163	1.00	28.74
5	OTA 5	M 558		THR	187	39.997	74.456	-6.203	1.00	28.71
	ATO			GLY	183	39.561 38.908	74.430	-5.619	1.00	0.00
	ATO			GLY	188	40.932	74.607	-5.753	1.00	25.69
	ATO			GLY	188 188	41.872	73.681	-6.483	1.00	27.48
_	ATO		_	GLY	188	42.983	74.024	-6.871	1.00	28.61
6				GLY MET	189	41.491	72.438	-6.720	1.00	27.34
	ATC			MET	189			-6.415	1.00	0.00
	ATC			MET	189			-7.426	1.00	25.62
	ATC			MET	189	_		-7.218	1.00	31.58
	ATC	OM 56	م ،	1-10-1						

	ATOM	568	CG	MET	189	42.445	69.542	-5.843	1.00	37.74
	ATOM	569	SD	MET	189	44.201	69.754	-5.409	1.00	37.50
	ATOM	570	CE	MET	189	44.983	68.533	-6.417	1.00	38.22
5	MOTA MOTA	571 572	C	MET	189	42.477	71.893	-8.893	00.1	24.18
,	ATOM	573	O N	MET THR	189	43.559	71.814	-9.460 0.670	1.00	27.99
	ATOM	574	H	THR	190 190	41.385	72.251	-9.570	1.00	24.30
	ATOM	575	CA	THR	190	40.494 41.446	72.204 72.677	-9.158	00.1	0.00
	ATOM	576	CB	THR	190	40.030	73.023	-10.959 -11.453	1.00 1.00	23.16
10	ATOM	577	OG1	THR	190	39.259	71.861	-11.433 -11.228	1.00	23.46 23.54
	ATOM	578	HGI	THR	190	38.355	72.025	-11.500	00.1	0.00
	MOTA	579	CGŹ	THR	190	39.922	73.321	-12.943	1.00	19.04
	MOTA	580	C	THR	190	42.363	73.878	-11.143	1.00	25.40
	MOTA	581	0	THR	190	43.255	73.913	-11.989	1.00	25.28
15	ATOM	582	N	MET	191	42.207	74.935	-10.353	1.00	25.85
	MOTA	583	H	MET	191	41.531	74.956	-9 .640	1.00	0.00
	MOTA	584	CA	MET	191	43.092	76.058	-10.563	1.00	26.80
	MOTA	585	CB	MET	191	42.618	77.263	<i>-9.7</i> 31	1.00	31.82
	ATOM	5 86	CG	MET	191	41.203	77.811	-10.021	1.00	30.18
2 0	MOTA	<i>5</i> 87	SD	MET	191	40.720	77.816	-11.767	1.00	42.05
	MOTA	<i>5</i> 88	CE	MET	191	42.005	78.898	-12.322	1.00	40.61
	MOTA	589	C	MET	191	44.534	75.727	-10.227	1.00	26.94
	ATOM	590	0	MET	191	45.439	76.102	-10.949	1.00	28.00
25	MOTA	591	N	LEU	192	44.841	75.015	-9.139	1.00	25.81
25	MOTA MOTA	592 593	H	LEU	192	44.141	74.785	-8.487	1.00	0.00
	ATOM	593 594	CA CB	LEU LEU	192	46.207	74.599	-8.872	1.00	22.18
	ATOM	595	CG	LEU	192 192	46.226 47.592	73.695 73.125	-7.637 -7.246	1.00 1.00	19.29 25.33
	ATOM	596	CD1	LEU	192	48.470	74.276	-7.240 -6.797	1.00	25.29
30	ATOM	597	CD2	LEU	192	47.450	72.049	-6.153	1.00	25.35
	ATOM	598	C	LEU	192	46.798	73.872	-10.049	1.00	19.90
	MOTA	599	0	LEU	192	47.871	74.187	-10.546	1.00	19.35
	ATOM	600	N	LEU	193	46.115	72.857	-10,554	1.00	20.99
	MOTA	601	H	LEU	193	45.208	72.654	-10.237	1.00	0.00
35	MOTA	602	CA	LEU	193	46.743	72.049	-11.571	1.00	21.52
	MOTA	603	CB	LEU	193	45.920	70.802	-11.849	1.00	21.31
	ATOM	604	CG	LEU	193	46.081	69.749	-10.749	1.00	23.69
	ATOM	605	CDI	LEU	193	45.125	68.587	-11.000	1.00	26.09
40	ATOM ATOM	606 607	CD2	LEU	193	47.509	69.236	-10.751	1.00	24.27
±0	ATOM	608	C 0	Leu Leu	193 193	46.921	72.815	-12.844	1.00	25.18
	ATOM	609	N	GLN	193	47.962 45.909	72,768 73,565	-13.484 -13.262	1.00 1.00	25.97 27.74
	ATOM	610	H	GLN	194	45.038	73.515	-13.202 -12.80 9	1.00	0.00
	ATOM	611	CA	. GLN	194	46.071	74.455	-14.391	1.00	29.82
45	ATOM	612	СВ	GLN	194	44.769	75.244	-14.530	1.00	28.29
	ATOM	613	CG	GLN	194	44.400	75.460	-15.983	1.00	33.28
	ATOM	614	CD	GLN	194	42.994	75.954	-16.101	1.00	34.79
	ATOM	615	OEI	GLN	194	42.085	75.270	-16.546	1.00	33.62
	ATOM	616	NE2	GLN	. 194	42.733	77.202	-15.779	1.00	36.58
50	ATOM	617	HE21	GLN	194	43.468	77.783	-15.495	1.00	0.00
	ATOM	618	HE22	GLN	194	41.799	77.487	-15.861	1.00	0.00
	MOTA	619	C	GLN	194	47.281	75.347	-14.146	1.00	31.33
	MOTA	620	0	GLN	194	48.177	75.481	-14.964	1.00	31.54
55	MOTA	621	N	ASN	195	47.356	75.986	-12.990	1.00	33.01
25	ATOM ATOM	622 623	H	ASN	195	46.613	75.928	-12.354	1.00	0.00
	ATOM	624	CA CB	asn Asn	195	48.530	76.762	-12.606	1.00	30.77
	ATOM	625	CG	ASN	195	48.460	77.239	-11.151	1.00	34.88
	ATOM	626	OD1	ASN NZA	195 195	47.691 47.679	78.525 79.452	-10.957 -11.753	1.00 1.00	38.35 43.57
60	MOTA	627	ND2	ASN	195	47.015	78.668	-11.733 -9.836	1.00	36.28
	ATOM	628	HD21	ASN	195	47.013	77.925	-9.206	1.00	0.00
	ATOM	629	HD22	ASN	195	46.506	79.497	-9.715	1.00	0.00
	MOTA	630	C	ASN	195	49.860	76.066	-12.719	1.00	28.51
	ATOM	631	ō	ASN	195	50.871	76.696	-12.949	1.00	26.10

	ATOM	632	N	LEU	196	49.928	74.758	-12.542	1.00	27.38
	ATOM	633	H	LEU	196	49.112	74.264	-12.313	1.00	0.00
	ATOM	634	CA	LEU	196	51.193	74.05l	-12.683	1.00	25.33
	MOTA	635	CB	LEU	196	51.179	72.795	-11.806	1.00	24.19 23.30
5	MOTA	636	CG	LEU	196	51.188	73.150	-10.32 6	1.00	24.60
	ATOM	637	CDI	LEU	196	50.972	71.914	-9.481 -9.985	1.00	31.63
	MOTA	638	CD2	LEU	196	52.519	73.794 73.661	-9.963 -14.123	1.00	26.23
	MOTA	639	C	LEU LEU	196 196	51.433 52.442	73.001	-14.503	1.00	24.03
10	MOTA MOTA	640 641	O N	GLY	197	50.478	73.970	-14.993	1.00	27.17
10	MOTA	642	H	GLY	197	49.631	74.363	-14.695	1.00	0.00
	ATOM	643	CA.	GLY	197	50.704	73.779	-16.401	1.00	25.75
	ATOM	644	C	GLY	197	49.860	72.707	-17.015	1.00	26.04
	ATOM	645	٥	GLY	197	50.142	72.320	-18.140	1.00	27.75
15	MOTA	646	N	TYR	198	48.830	72.188	-16.360	1.00	24.74
	ATOM	647	H	TYR	198	48.531	72.597	-15.518	1.00	0.00 24.17
	MOTA	648	CA	TYR	198	48.125	71.016	-16.872	1.00 1.00	24.17
	MOTA	649	СВ	TYR	198	47.870	69.968	-15.744 -15.195	1.00	17.56
	MOTA	650	CG	TYR	198	49.145 49.919	69.319 69.969	-14.251	1.00	17.53
20	ATOM	651	CD1	TYR TYR	198 198	51.123	69.433	-13.835	1.00	19.28
	MOTA MOTA	652 653	CD2	TYR	198	49.572	58.117	-15.711	1.00	16.54
	MOTA	654	CES	TYR	198	50.771	67.572	-15.298	1.00	17.99
	ATOM	655	cz	TYR	198	51.567	68.265	-14.3 9 9	1.00	22.77
25	MOTA	656	OH	TYR	198	52.806	67.774	-14.022	1.00	26.91
	ATOM	657	HH	TYR	198	53.179	68.383	-13.376	1.00	0.00
	ATOM	658	C	TYR	198	46.794	71.385	-17.465	1.00	25.30
	MOTA	659	0	TYR	198	46.136	72.288	-16.959	1.00	28.39 27.93
	ATOM	660	N	SER	199	46.353	70.714	-18.533 -19.092	1.00	0.00
30	ATOM	661	H	SER	199	46.951 44.953	70.163 70.775	-19.092	1.00	29.13
	ATOM	662	CA CB	SER SER	199 199	44.933 44.776	70.400	-20.402	1.00	31.52
	MOTA ATOM	663 664	OG	SER	199	45.617	71.216	-21.197	1.00	44.44
	MOTA	665	HG	SER	199	45.356	72.131	-21.040	1.00	0.00
35	ATOM	666	c	SER	199	44.163	69.810	-18.077	1.00	25.88
	ATOM	667	ō	SER	199	44,489	68.636	-17.967	1.00	28.99
	ATOM	668	N	VAL	200	43.095	70.325	-17.471	1.00	22.17
	ATOM	669	H	VAL	200	42.823	71.240	-17.708	1.00 1.00	0.00 21.46
	ATOM	670	CA	VAL	200	42.321	69.612	-16.481	1.00	17.18
40	MOTA		CB	VAL	200	42.090	70.526 69.691	-15.284 -14.142	1.00	19.27
	MOTA		CG1 CG2	VAL VAL	200 200	41.528 43.399	71.222	-14.885	1.00	17.36
	MOTA		C	VAL	200	40,978	69.116	-16.979	1.00	25.60
	MOTA MOTA		ō	VAL	200	40.073	69.889	-17.248	1.00	26.95
45	ATOM		Ň	ASP	201	40.787	67.815	-17.115	1.00	25.10
	ATOM		н	ASP	201	41.548	67.209	-17.020	1.00	0.00
	ATOM		CA	ASP	201	39.459	67.259	-17.292	1.00	26.46
	ATOM	679	CB	ASP	201	39.501	65.852	-17.808	1.00	32.29
	ATOM	680	CG	ASP	201	39.387	65.818	-19.271	1.00	40.02 53.36
50	ATOM		ODI	ASP	201	38.413	65.246	-19.737	1.00 1.00	48.96
	ATOM		OD2	ASP	201	40.251	66.358	-19,949	1.00	28.68
	ATOM		C	ASP	201	38.786	67.159 66.593	-15.940 -15.030	1.00	25.09
	ATOM		0	ASP	201 202	39.402 37.561	67.665	-15.755	1.00	26.15
	ATOM		N	VAL VAL	202	37.092	68.158	-16.463	1.00	0.00
55	MOTA MOTA		H CA	VAL	202	36.820	67.390	-14.543	1.00	29.60
			CB	VAL	202	36.274	68.690	-13.955	- 1.00	29.43
	KOTA KOTA		CGI	VAL	202	35.461	68,387	-12.696	1.00	26.97
	ATON		CG2	VAL	202	37.434	69.634	-13.649	1.00	26,15
60			c	VAL	202	35.669	66.447	-14.865	1.00	33.37
~ ~	MOTA		ō	YAL	202	34.893	66.731	-15.763	1.00	36.29
	ATO		N	LYS	203	35.496	65.309	-14.183	1.00	33.12
	ATO	M 694	н	LYS	203	36.163	65.050	-13.508	1.00	0.00 28.52
	ATO		CA	LYS	203	34.334	64.448	-14.372	1.00	20.34

- 53 -

	ATOM	696	CB	LYS	203	34.749	63.050	-14.826	1.00	28.62
	ATOM	697	CG	LYS	203	35.590	62.990	-16.088	1.00	32.11
	MOTA	698	CD	LYS	203	34.746	63.437	-17.266	1.00	40.39
_	ATOM	699	CE	LYS	203	35.531	63.532	-18.574	1.00	50.57
5	MOTA	700	NZ	LYS	203	34.651	64.012	-19.625	1.00	59.39
	MOTA	701	HZI	LYS	203	34.274	64.946	-19.364	1.00	0.00
	ATOM	702 703	HZ2	LYS	203	33.854	63.352	-19.738	1.00	0.00
	ATOM	703 704	HZ3 C	LYS	203	35.168	64.088	-20.524	1.00	0.00
10	ATOM	705	Ö	LYS LYS	203	33.596	64.313	-13.046	1.00	29.68
	ATOM	706	N	LYS	203	34.220	64.117	-12.016	1.00	30.93
	ATOM	707	н.	LYS	204 204	32.268	64.402	-12.954	1.00	32.55
	ATOM	708	CA	LYS	204	31.711 31.610	64.391 64.417	-13.762	1.00	0.00
	ATOM	709	СВ	LYS	204	30.827	65.701	-11.655	1.00	34.17
15	ATOM	710	CG	LYS	204	31.646	66.950	-11.363 -11.553	1.00 1.00	32.07 37.18
	ATOM	711	CD	LYS	204	30.858	68.196	-11.188	1.00	44.8
	ATOM	712	CE	LYS	204	31.660	69.462	·11.544	1.00	54.50
	ATOM	713	NZ	LYS	204	30.997	70.674	-11.086	1.00	60.66
	ATOM	714	HZI	LYS	204	30.884	70.631	-10.052	1.00	0.00
20	MOTA	715	HZ2	LYS	204	30.058	70.742	-11.528	1.00	0.00
-	MOTA	.716	HZ3	LYS	204	31.568	71.505	-11.339	1.00	0.00
	ATOM	717	¢	LYS	204	30.603	63.311	-11.602	1.00	32.66
	ATOM	718	0	LYS	204	30.073	62.894	-12.625	1.00	33.79
25	ATOM	719	N	ASN	205	30.378	62.86 6	-10.365	1.00	31.17
23	MOTA MOTA	720	H	ASN	205	30.983	63.186	·9.659	1.00	0.00
	ATOM	721 722	CA CB	ASN	205	29.340	61.902	-10.003	1.00	29.81
	ATOM	723	CG	ASN ASN	205	27.956	62.531	-10.122	1.00	25.29
	ATOM	724	ODI	ASN	205 205	27.915	63.623	-9.103	1.00	26.48
30	ATOM	725	ND2	ASN	205	28.416 27.363	63.536 64.768	-7.988	1.00	29.97
	ATOM	726	HD21	ASN	205	27.014	64.830	-9.456 -10.369	1.00 1.00	29.11 0.00
	ATOM	727	HD22	ASN	205	27.318	65.493	-10.30 3 -8.798	1.00	0.00
	ATOM	728	С	ASN	205	29.323	60.626	-10.792	1.00	30.34
	ATOM	729	0	ASN	205	28.356	60.317	-11.453	1.00	32.35
35	MOTA	730	N	LEU	206	30.354	59.799	-10.793	1.00	31.19
	MOTA	731	H	LEU	206	31.092	59.947	-10.163	1,00	0.00
	ATOM	732	CA	LEU	206	30.352	58.667	-11.6 9 8	1.00	28.61
	ATOM	733	CB	LEU	206	31.657	58.587	-12.498	1.00	29.20
40	ATOM	734	CG.	LEU	206	32.070	5 9.852	-13.281	1.00	29.45
40	MOTA MOTA	735 736	CD1 CD2	LEU	206	33.375	59.557	-14.009	1.00	30.38
	ATOM	737	C	LEU LEU	206	31.010	60.255	-14.297	1.00	33.07
	ATOM	738	ò	LEU	206 206	30.196 30.211	57.402	-10.933	1.00	30.76
	ATOM	739	N	THR	207	30.211	57.314 56.331	-9.715 -11.664	1.00	35.60
45	ATOM	740	H	THR	207	29.939	56.423	-12.631	1.00 1.00	32.87 0.00
	ATOM	741	CA	THR	207	30.012	55.027	-11.058	1.00	36.34
	ATOM	742	CB	THR	207	28.851	54.344	-11.773	1.00	36.91
	ATOM	743	OG1	THR	207	27.728	54.701	-10.986	1.00	42.24
	MOTA	744	HGI	THR	207	26.926	54.355	-11.394	1.00	0.00
50	ATOM	745	CG2	THR	207	28.942	52.841	-11.905	1.00	42.72
	ATOM	746	C	THR	207	31.381	54.358	-11.219	1.00	36.64
	MOTA	747	0	THR	207	32.157	54.742	-12.079	1.00	39.32
	ATOM	748	N	ALA	208	31.747	53.344	-10.429	1.00	36.10
55	ATOM ATOM	749 750	H	ALA	208	31.129	53.028	-9.745	1.00	0.00
55	ATOM	751	CA CB	ALA ALA	208	33.047	52.699	-10.537	1.00	34.66
	ATOM	752	C	ALA ALA	208 208	33.042	\$1.453	-9.643	1.00	32.92
	ATOM	753	ō	ALA	208	33.402	52.325 52.467	-11.959	1.00	36.07
	ATOM	754	й	SER	208	34.525 32.419	52.467 51.840	-12.425 -12.707	1.00 1.00	38.58
60	ATOM	755	н	SER	209	31.516	51.718	-12.707	1.00	38.40
	ATOM	756	CA	SER	209	32.623	51.470	-12.349 -14.0 9 6	1.00	0.00 18.81
	ATOM	757	CB	SER	209	31.471	50.593	-14.573	1.00	45.07
	ATOM	758	OG	SER	209	30.259	51.040	-13.965	1.00	56.06
	ATOM	759	HG	SER	209	29.534	50,500	-14.307	1.00	0.00

										27.04
	MOTA	760	С	SER	209	32.724	52.676	-14.978	1.00 1.00	35.05 36.86
	MOTA	761	0	SER	209	33.571	52.733	-15.857 -14.779	1.00	33.45
	MOTA	762	N	ASP	210	31.877	53.687 53.563	-14.179	1.00	0.00
-	MOTA	763	H	ASP	210	31.138 32.061	54.930	-15.516	1.00	36.67
5	MOTA	764	CA CB	ASP ASP	210 210	31.103	56.049	-15.088	1.00	43.07
	MOTA MOTA	765 766	CG	ASP	210	29.602	55.780	-15.228	1.00	51.26
	ATOM	767	ODI	ASP	210	29.209	54.945	-16.060	1.00	52.07
	ATOM	768	OD2	ASP	210	28.830	56.441	-14.507	1.00	50.26
10	ATOM	769	C	ASP	210	33.469	55.432	-15.219	1.00	36.66
	MOTA	770	0	ASP	210	34.155	55.935	-16.096	1.00	34.14
	MOTA	771	N.	MET	211	33.944	55.299	-13.965	1.00 1.00	36.25 0.00
	ATOM	772	H	MET	211	33.371	54.919	-13.268 -13.629	1.00	36.30
٦.	ATOM	773	CA	MET	211	35.310 35.574	55.693 55.528	-12.130	1.00	32.55
15	MOTA	774 775	CB CG	MET MET	211 211	34.914	56.611	-11.271	1.00	31.33
	ATOM ATOM	776	SD	MET	211	35.182	56.360	-9,493	1.00	38.31
	ATOM.	777	CE	MET	211	34.219	57.715	-8.954	1.00	38.56
	ATOM	778	c	MET	211	36.310	54.866	-14.404	1.00	33.55
20	ATOM	779	0	MET	211	37.247	55.382	-14. 999	1.00	35.45
•	MOTA	780	N	THR	212	36.140	53.551	-14.436	1.00 1.00	30.54 0.00
	MOTA	781	H	THR	212	35.445	53.118 52.704	-13.893 -15.242	1.00	31.66
	ATOM	782	CA	THR	212	36.997 36.443	51.289	-15.170	1.00	30.49
25	MOTA	783 7 84	CB OG1	THR THR	212 212	36.580	50.901	-13.805	1.00	36.43
25	MOTA MOTA	785	HG1	THR	212	37.491	50.999	-13.515	1.00	0.00
	ATOM	786	CG2	THR	212	37.128	50.325	-16.127	1.00	30.70
	ATOM	787	c	THR	212	37.122	53.142	-16.681	1.00	30.78
	MOTA	788	0	THR	212	38.195	53.275	-17.241	1.00	30.06
30	ATOM	789	N	THR	213	35.973	53.375	-17.297	1.00 1.00	34.48 0.00
	MOTA	790	H	THR	213	35.126	53.245 53.706	-16.813 -18.681	1.00	33.70
	ATOM	791	CA	THR	213 213	35.869 34.354	53.796 53.892	-19.044	1.00	38.71
	KOTA	792 793	CB OG1	THR THR	213	33.818	52.581	-18.856	1.00	42.49
35	MOTA MOTA	793 794	HG1	THR	213	34.273	51.966	-19.436	1.00	0.00
22	ATOM	795	CG2	THR	213	34.086	54,420	-20.467	1.00	40.74
	ATOM		c	THR	213	36.571	55.122	-18.900	1.00	31.43
	ATOM	797	0	THR	213	37.304	55.298	-19.865	1.00	30.44 29.82
	MOTA		N	GLU	214	36.356	56.097	-18.006 -17.279	1.00 1.00	0.00
40	MOTA		H	GLU	214	35.706	55.963 57.3 51	-18.103	1.00	29.65
	ATOM		CA	GLU GLU	214 214	37.082 36.673	58.355	-17.025	1.00	33.21
	MOTA MOTA		CB CG	GLU	214	35.275	58.972	-17.161	1.00	39.44
	ATOM		CD	GLU	214	34.998	59.508	-18.559	1.00	44.47
45			OE1	GLU	214	35.754	60.353	-19.052	1.00	46.83
	ATOM		OE2	GLU	214	34.015	5 9.057	-19.154	1.00	48.55 26.00
	ATOM	806	C	GLU	214	38.561	57.147	-17.960	1.00 1.00	23.58
	ATOM		0	GLU	214	39.348	5 7.759 5 6.276	-18.665 -17.045	1.00	26.22
	ATOM		N	LEU	215 · · 215	38.976 38.327	55.881	-16.422	1.00	0.00
50			H	LEU LEU	215	40.392	55.922	-16.942	1.00	30.25
	MOTA		CA CB	LEU	215	40.670	54.929	-15.786	1.00	28.64
	ATON		CG	LEU	215	40.608	55.547	-14.386	1.00	31.22
	ATON		CDI	LEU	215	40.822	54.470	-13.329	1.00	30.89
55			CD2	LEU	215	41.648	56.656	-14.292	1.00	24.41
	ATON		С	LEU	215	40.966	55.300	-18.190	1.00	29.50 32.89
	KOTA		0	LEU	215	41.998	55.738	-18.686	1.00 1.00	27.46
	ATO		N	GLU	216	40.334	54.261 53.884	-18.740 -18.282	1.00	0.00
	ATO:		H	GLU	216	39.555 40.802	53.884 53.695	-19.991	1.00	30.27
60			CA	GLU GLU	216 216	39.922	52.537	-20.463	1.00	38.22
	OTA		C8 C8	GLU	216	39.940	51.319	-19.517	1.00	54.67
	ATO		CD	GLU	216	39.089	50.173	-20.064	1.00	60.29
	ATO			GLU	216	39.535	49.546	-21.027	1.00	63.24

- 55 -

	ATOM	824	OE2	GLU	216	38.003	49.907	-19.525	1.00	60.46
	ATOM	825	С	GLU	216 .	40.823	54.724	-21.094	1.00	29.88
	ATOM	826	0	GLU	216	41.722	54.757	-21.926	00.1	27.84
_	MOTA	827	N ·	ALA	217	39.836	55.617	-21.144	1.00	27.72
5	ATOM	828	н	ALA	217	39.086	\$5.\$50	-20.516	1.00	0.00
	MOTA	829	CA	ALA	217	39.851	56.698	-22,121	1.00	27.12
	ATOM	830	CB	ALA	217	38.530	57.477	-22.045	1.00	27.01
	MOTA	831	.c	ALA	217	41.008	<i>5</i> 7.620	-21.855	1.00	26.62
10	MOTA MOTA	832	0	ALA	217	41.791	57.934	-22.740	1.00	26.18
10	ATOM	833 834	N H	PHE	218	41.179	58.083	-20.624	1.00	25.15
	ATOM	835	CA	PHE	218	40.554	57.809	-19.917	1.00	0.00
	MOTA	836	CB	PHE PHE	218	42.258	58.999	-20.319	1.00	23.19
	ATOM	837	CG	PHE	218 218	42.185	59.322	-18.823	1.00	20.84
15	ATOM	838	CDI	PHE	218	43.236 44.361	60.340	-18.445	1.00	25.38
	ATOM	839	CD2	PHE	218	43.036	59.929 61.686	-17.742 -18.721	00.1 00.1	24,12 23,15
	ATOM	840	CEI	PHE	218	45.234	60.889	-17.262	1.00	23.13
	ATOM	841	CE2	PHE	218	43.920	62.634	-18.228	1.00	23.97
	MOTA	842	CZ	PHE	218	45.005	62.234	-17.484	1.00	21.26
20	ATOM	843	C	PHE	218	43.603	58.427	-20.701	1.00	25.14
•	ATOM	844	0	PHE	218	44.495	59.110	-21.209	1.00	29.00
	MOTA	845	N	ALA	219	43.772	57.138	-20.456	1.00	23.61
	ATUM	846	H	ALA	219	43.083	56.655	-19.953	1.00	0.00
	ATOM	847	CA	ALA	219	44.971	56.429	-20.858	1.00	24,15
25	ATOM	848	CB	ALA	219	44.884	54.9 79	-20.375	1.00	24.62
	ATOM	849	C	ALA	219	45.224	56.435	-22.351	1.00	30.25
	MOTA MOTA	850 851	и О	ALA	219	46.335	56.291	-22.834	1.00	30.75
	ATOM	852	H	HIS HIS	220	44.155	56.608	-23.131	1.00	31.80
30	ATOM	853	CA	HIS	220 220	43.276 44.260	56.759	-22.722	1.00	0.00
7.	ATOM	854	CB	HIS	220	43.030	56.691 56.108	-24.578 -25.287	1.00 1.00	32.54
	MOTA	855	CG	HIS	220	43.153	54.627	-25.144	1.00	33.65 38.22
	ATOM	856	CD2	HIS	220	43.712	53.790	-26.069	1.00	40.87
	MOTA	857	ND1	HIS	220	42.884	53.894	-24.069	1.00	2.77
35	ATOM	858	HD1	HIS	220	42.460	54.202	-23.244	1.00	0.00
	MOTA	859	CEI	HIS	220	43.283	52.673	-24.281	1,00	41.06
	ATOM	860	NE2	HIS	220	43.788	52.624	-25.483	1.00	39.69
	ATOM	861	HE2	HIS	220	44.189	51.815	-25.855	1.00	0.00
4.0	ATOM	862	C	HIS	220	44.431	58.066	-25.139	1.00	31.27
40	MOTA	863	0	HIS	220	44.482	58.222	-26.346	1.00	34.54
	MOTA MOTA	864 865	N H	ARG	221	44.519	59.129	-24.354	1.00	31.23
	ATOM	866	CA	ARG ARG	2 21	44.453	59.018	-23.381	1.00	0.00
	ATOM	867	CB	ARG	221 221	44.684 44.496	60.441 61.460	-24.935	1.00	25.20
45	ATOM	868	CG	ARG	221	43.089	61.433	-23.847 -23.298	1.00 1.00	23.87 24.28
	ATOM	869	CD	ARG	221	42.164	62.262	-24.150	1.00	24.28 24.77
	ATOM	870	NE	ARG	221	42.527	63.666	-24.117	1.00	30.34
	ATOM	871	HE	ARG	221	43.163	64.003	-24.780	1.00	0.00
	ATOM	872	CZ	ARG	221	42.025	64.537	-23.214	1.00	36.39
50	ATOM	873	NHI	ARG	221	41.323	64.137	-22.120	1.00	37.62
	ATOM	874	HH 11	ARG	2 21	41.185	63.163	-21.950	1.00	0.00
	ATOM	875	HH 12	ARG	221	40.981	64.814	-21.467	1.00	0.00
	MOTA	876	NH2	ARG	221	42.162	65.884	-23.412	1.00	36.36
E E	ATOM	877	HH21	ARG	221	42.631	66.223	-24.228	1.00	0.00
55	ATOM	878	HH22	ARG	221	41.797	66.529	-22.740	1.00	0.00
	ATOM	879	C	ARG	221	46.030	60.643	-25.605	1.00	27.94
	ATOM ATOM	880 881	0 N	ARG	221	47.097	60.475	-25.015	1.00	27.32
	ATOM	882 882	N CD	PRO PRO	222	46.015	61.030	-26.842	1.00	30.14
60	ATOM	883	CA	PRO	222 222	44.795	61.277	-27.621	1.00	28.66
~ •	MOTA	884	CB	PRO	222	47.204 46.670	61.317	-27.635 -28.976	1.00	26.80
	ATOM	885	CG	PRO	222	45.279	61.799 62.298	-28.976 -28.634	1.00 1.00	27.03 26.03
	ATOM	886	č	PRO	222	48.125	62.326	-26.990	1.00	26.07 28.33
	ATOM	887	ŏ	PRO	222	49.329	62.305	-20.990 -27.184	1.00	26.33 34.07
			-				02.303	-21.104	1.00	3−7. 07

		000	N	•	LU	223	47.627	63.254	-26.189	1.00	25.34
	ATOM	888 889	H		SLU	223	46.669	63.259	-26.005	1.00	0.00
	MOTA	890	CA		3LU	223	48.500	64.265	-25.620	1.00	28.40
	ATOM	891	CB		GLU	223	47.671	65.306	-24.887	1.00	27.51 32.04
5	ATOM	892	CG		JLU	223	46.658	66.030	-25.769	1.00 1.00	34.72
	ATOM	893	CD	- (GLU	223	45.293	65.381	-25.726	1.00	35.25
	ATOM	894	OE1		GLU	223	44.328	66.114	-25.534	00.1	38.35
	ATOM	895	OE2		GLU	223	45.177	64.168	-25.880 -24.663	1.00	28.17
	ATOM	896	C	1	GLU	223	49.569	63.758	-24.003 -24.461	1.00	28.18
10	MOTA	897	0		GLU	223	50.606	64.371	-24.034	1.00	26.29
-	MOTA	898	N		HIS	224	49,343	62.599 62.108	-24,230	1.00	0.00
	MOTA	899	H.		HIS	224	48.522 50.346	62.037	-23.139	1.00	25.11
	MOTA	900	CA		HIS	224	49.960	60.670	-22.601	1.00	21.67
	ATOM	901	CB		HIS	224 224	48.774	60.779	-21.676	1.00	28.93
15	MOTA	902	CG		HIS	224 224	47.721	59.901	-21.673	1.00	29.23
	ATOM	903	CD:		HIS	224	48.546	61.704	-20.722	1.00	26.59
	MOTA	904	מא		HIS HIS	224	49.236	62.337	-20.408	1.00	0.00
	MOTA	905	HD		HIS	224	47.382	61.423	-20.177	1.00	31.44
	ATOM	906	CE NE		HIS	224	46.889	60.352	-20.773	1.00	33.43
2,0	MOTA	907 908	HE		HIS	224	46.004	59.939	-20.625	1.00	0.00
	MOTA MOTA	909	C	-	HIS	224	51.646	61.845	-23.859	1.00	26.89
	MOTA	910	ŏ		HIS	224	52,700	62.029	-23.281	1.00	30.62 30.81
	MOTA		N		LYS	225	51.608	61.474	-25,138	1.00	0.00
25	MOTA		н		LYS	225	50.750	61.290	-25.584	1.00	32.77
25	ATOM		CA		LYS	225	52,826	61.359	-25.937	1.00 1.00	39.91
	ATOM		CE		LYS	225	52.5 03	60.985	-27,373	1.00	55.31
	ATOM		CC	;	LYS	225	52.645	59.492	-27.564 -29.024	1.00	67.66
	ATOM		CI)	LYS	225	52.548	59.057	-29.181	1.00	75.00
30	ATOM	917			LYS	225	52.571	57.524 56.917	-28.699	1.00	78.51
	ATOM				LYS	225	51.337 50.529	57.292	-29.236	1,00	0.00
	ATOM				LYS	225 225	51.211	57.140	-27.691	1.00	0.00
	ATOM				LYS	225	51.387	55.887	-28.827	1.00	0.00
	ATOM			Z3	LYS LYS	225	53.638	62.627	-25.985	1.00	30.17
35	ATOM				LYS	225	54.854	62.599	-26.051	1.00	32.06
	ATOM				THR	226	53.015	63.792	-25.975	1.00	29.27
	ATON				THR	226	52.045	63.869	-26.061	1.00	0.00 29.35
	AOTA AOTA			A	THR	226	53.803	64.992	-25.854	1.00	28.73
40				B	THR	226	53.293	66.018	-26.896	1.00	27.05
40	ATO		•	G1	THR	226	51.944	65.709	-27.203	1.00 1.00	0.00
	ATO			(G1	THR	226	51.452	66.009	-26.422 -28.180	1.00	31.09
	ATO		0 0	:G2	THR	226	54.127	65.983	-24.441	1.00	30.98
	ATO		1 (:	THR	226	53.770	65.546 66.756	-24.226	1.00	34.36
45	ATO	M 93			THR	226	53.809	64.702	-23.408	1.00	27.84
	ATO	M 93			SER	227	53.685 53.657	63.727	-23.528	1.00	0.00
	OTA			i.	SER	227	53.733	65.196	-22.055	1.00	25.31
	OTA		-	CA	SER	227 227	52.412	64.917	-21.322	1.00	23.69
	OTA			CB	SER	. 227	52.319	65.533	-20.021	1.00	29.36
50				DG C	ser ser	227	51.584	65.178	-19.486	1.00	0.00
	ATO			HG	SER	227	54.872	64.535	-21.301	1.00	27.92
	ATC			c o	SER	227	55.409	63.498	-21.661	1.00	22.67
	ATC			N	ASP	228	55,246	65.201	-20.199	1.00	28.65
-	ATC 5 ATC			H	ASP	228	54.753	66.020	-19.977	1.00	0.00
5	OTA		43	CA	ASP	228	55,270	64.715	-19.270	1.00	28.62 21.37
	ATO		44	CB	ASP	228	57.305	65.848	-18.973	1.00	
	TA		145	CG	ASP	228			-18.438	1.00	23.40 25.53
	AT(746	ODI	ASP	228	57.311		-18.499	1.00	24.44
6	O AT		947	OD2	ASP	228			-17.976		29.79
0	AT		948	C	ASP	228					33.42
	TA		949	0	ASP	228					30.06
			950	N	SER	229					0.00
			951	H	SEX	229	53.756	64.595	-10.213		J.54

- 57 -

	MOTA		CA	SER	229	53.905	63.708	-16.336	1.00	24.85
	ATOM		CB	SER	229	54.314	64.599	-15.182	1.00	21.77
	MOTA MOTA		OG	SER	229	54.171	65.969	-15.547	1.00	21.06
5	ATOM		HG C	SER	229	54.272	66.564	-14.792	1.00	0.00
•	ATOM		ŏ	SER SER	229	52.413	63.747	-16.437	1.00	23.31
	ATOM		N	THR	229 230	51.900	64.273	-17.423	1.00	23.18
	ATOM	959	H	THR	230	51.720 52.164	63.201 62.618	-15.435	1.00	24.93
	ATOM	960	CA	THR	230	50.303	63.473	-14.780 -15.239	1.00 1.00	0.00 22.17
10	MOTA	961	CB	THR	230	49.501	62.410	-16.052	1.00	18.90
	MOTA	962	OĞİ	THR	230	48.158	62.903	-16.198	1.00	23.08
	ATOM	963	HG1	THR	230	48.158	63.784	-16.611	1.00	0.00
	MOTA MOTA	964 965	CG2	THR	230	49.485	61.030	-15.391	1.00	17.19
15	ATOM	966	C O	THR THR	230	49.933	63.471	-13.739	1.00	22.19
	ATOM	967	N	PHE	230 231	50.683	63.015	-12.887	1.00	19.95
	ATOM	968	н	PHE	231	48.751 48.196	63.999 64.345	-13.412	1.00	23.01
	ATOM	969	CA	PHE	231	48.168	64.022	-14.142 -12.081	1.00 1.00	0.00
	ATOM	970	CB	PHE	231	47.955	65.441	-11.549	1.00	22.98 22.01
20	ATOM	971	CG	PHE	231	49.190	65.978	-10.873	1.00	23.66
	ATOM	972	CDI	PHE	231	49.220	66.071	-9.486	1.00	22.17
	ATOM	973	CD2	PHE	231	50.273	66.392	-11.635	1.00	21.19
	ATOM	974	CEI	PHE	231	50.361	66.561	-8.861	1.00	23.83
25	MOTA	975 976	CE2	PHE	231	51.391	66.900	-11.010	1.00	18.70
2,5	ATOM	977	CZ C	PHE PHE	231	51.442	66.965	-9.620	1.00	22.93
	ATOM	978	õ	PHE	231 231	46.792 46.067	63.409	-12.160	1.00	21.75
	ATOM	979	N	LEU	232	46.343	63.809 62.479	-13.062 -11.312	1.00 1.00	21.91 20.85
	ATOM	980	H	LEU	232	46.949	62.069	-10.654	1.00	0.00
30	ATOM	981	CV	LEU	232	44.915	62.139	-11.293	1.00	19.87
	ATOM	982	CB	LEU	232	44.627	60.672	-11.588	1.00	23.90
	ATOM ATOM	983 984	CG	LEU	232	45.050	60.289	-12.995	1.00	30.22
	ATOM	985	CD1 CD2	LEU	232	46.486	59.760	-12.956	1.00	32.80
35	ATOM	986	CD2	LEU LEU	232	44.089	59.240	-13.555	1.00	32.58
	ATOM	987	ō	LEU	232 232	44.338 44.980	62.387 62.002	-9.941	00.1	18.92
	MOTA	988	N	VAL	233	43.174	62.998	-8.979 -9.754	1.00	22.64 20.69
	ATOM	989	H	VAL	233	42.629	63.238	-10.535	1.00	0.00
4.0	ATOM	990	CA	VAL	233	42.664	63.290	-8.423	1.00	22.20
40	ATOM	991	CB	VAL	233	42.519	64.812	-8.180	1.00	21.39
	ATOM	992	CGI	VAL	233	42.185	65.090	-6.706	1.00	20.83
	MOTA MOTA	993 994	c c	VAL	233	43.826	65.525	-8.539	1.00	27.21
	MOTA	995	ò	VAL VAL	233 233	41.291	62.645	-8.264	1.00	27.10
45	ATOM	996	N	PHE	233	40.429 41.049	62.803 61.906	-9.122 -7.176	1.00	24.87
	ATOM	997	Н	PHE	234	41.749	61.801	-6.495	1.00 1.00	24.72 0.00
	ATOM	998	CA	PHE	234	39.764	61.286	-6.938	1.00	23.25
	ATOM	999	CB	PHE	234	39.870	59.773	-6.767	1.00	20.01
E 0	MOTA	1000	CG	PHE	234	40.400	59.110	-8.005	1.00	21.08
50	ATOM	1001	CDI	PHE	234	39.525	58.555	-8.904	1.00	26.42
	MOTA MOTA		CD2	PHE	234	41.768	59.045	-8.222	1.00	25.79
	ATOM		CE1 CE2	PHE PHE	234	40.019	57.951	-10.046	1.00	29.30
	ATOM		CZ	PHE	234	42.262	58.445	-9.353	1.00	23.33
55	ATOM		C	PHE	234 234	41.379 39.259	57.910	-10.266	1.00	31.52
	ATOM	1007	ō	PHE	234	39.990	61.858 61.926	-5.650 -4.664	1.00 1.00	25.38 27.17
	MOTA	1008	N	MET	235	38.006	62.298	-5.583	1.00	26.22
	ATOM		H	MET	235	37.441	62.301	-6.384	1.00	0.00
C C	ATOM		CA	MET	235	37.451	62.780	4.326	1.00	25.64
60		1011	CB	MET	235	37.276	64.294	-4.345	1.00	24.45
		1012	00	MET	235	38.619	64.946	-4.681	1.00	30.66
	ATOM ATOM		SD CE	MET	235	38.601	66.742	-4.848	1.00	37.65
	ATOM		C	MET MET	23.5 23.5	38.003	66.982	-6.481	1.00	33.32
			-		233	36.120	62.112	-4.236	1.00	29.15

	ATOM 1016	0	MET	235	35.339	62.186	-5.175	1.00	29.75
	ATOM 1017	N	SER	236	35.808	61.432	-3.138	1.00	31.99
	ATOM 1018	H	SER	236	36.441	61.364	-2.382	1.00	0.00
	ATOM 1019	CA	SER	236	34.531	60.769	-2.995	1.00	32.91
5	ATOM 1020	CB	SER	236	34.412	59.596	-3.971	1.00	31.29
-	ATOM 1021	OG	SER	236	33.125	58. 9 96	-3.885	1.00	32.86
	ATOM 1022	HG	SER	236	32.46 3	59.641	-4.143	1.00	0.00
	ATOM 1023	,C	SER	236	34.462	60.246	-1.573	1.00	32.69 33.35
	ATOM 1024	0	SER	236	35.389	60.340	-0.778	1.00	33.10
10	ATOM 1025	N	ИIS	237	33.329	59.661	-1.203	1.00 1.00	0.00
	ATOM 1026	H	HIS	237	32.551	59.730	-1.783 -0.016	1.00	34.12
	ATOM 1027	CA	HIS	237	33.332	58.809 58.331	0.377	1.00	37.00
	ATOM 1028	CB	HIZ	237	31.882 31.079	59.375	1.141	1.00	42.87
	ATOM 1029	CG	HIS HIS	237 237	29.798	59.740	0.816	1.00	40.88
15	ATOM 1030	CD2 ND1	HIZ	237	31.457	60.128	2.188	1.00	41.19
	ATOM 1031 ATOM 1032	HD1	HIS	237	32.325	60.120	2.661	1.00	0.00
	ATOM 1032	CEI	HIS	237	30.460	60.934	2.446	1.00	43.14
	ATOM 1034	NE2	HIS	237	29.466	60.721	1.609	1.00	42.94
20	ATOM 1035	HE2	HIS	237	28.639	61.252	1.540	1.00	0.00
2,0	ATOM 1036	C	HIS	237	34.194	57.577	-0.321	1.00	33.55
	ATOM 1037	0	HIS	237	34.520	57.234	-1.462	1.00	32.97
	ATOM 1038	N	GLY	238	34.606	56.852	0.702	1.00	32.27 0.00
	ATOM 1039	H	GLY	238	34.429	57.073	1.646	1.00	31.94
25	ATOM 1040	CA	GLY	238	35.369	55.668	0.428	1.00	31.47
	ATOM 1041	С	GLY	238	35.217	54.774	1.609 2.707	1.00	29.62
	ATOM 1042	0	GLY	238	34.874	55.189	1.370	1.00	30.61
	ATOM 1043	N	ILE	239	35.475	53.512 53.236	0.474	1.00	0.00
	ATOM 1044	H	ILE	239	35.767	52.548	2.439	1.00	36.45
30	ATOM 1045	CA .	ILE	239 239	35.461 34.416	51.450	2.210	1.00	38.77
	ATOM 1046	CB	ILE ILE	239	33.067	52.124	2.180	1.00	42.42
	ATOM 1047	CG2 CG1	ILE ILE	239	34.645	50.683	0.928	1.00	42.77
	ATOM 1048 ATOM 1049	CDI	ILE	239	33.745	49.470	0.810	1.00	45.44
35	ATOM 1050	c .	ILE	239	36.822	51.930	2.435	1.00	37.10
35	ATOM 1050	Ö	ILE	239	37.640	52.175	1.572	1.00	37.25
	ATOM 1052	N	ARG	240	37.118	51.076	3.398	1.00	39.29
	ATOM 1053	H	ARG	240	36.434	50.847	4.063	1.00	0.00 42.79
	ATOM 1054	CA	ARG	240	38.428	50.462	3.465	1.00	45.84
40	ATOM 1055	CB	ARG	240	38.400	49.475	4.647	1.00	52.68
	ATOM 1056	CG	ARG	240	39.680	48.660	4.892	1.00 1.00	57.36
	ATOM 1057	CD	ARG	240	40.939	49.489	5.139 4.594	1.00	60.04
	ATOM 1058	NE	ARG	240	42,114	48.829 48.953	3.645	1.00	0.00
	ATOM 1059	HE	ARG	240	42.319	48.052	5.327	1.00	66.92
45			ARG	240	42.922 44.059	47,576	4,740	1.00	71.12
	ATOM 1061	NHI	ARG	240 240	44.259	47.811	3.789	1.00	0.00
	ATOM 1062		ARG	240	44.688	46.986	5.247	1.00	00.0
	ATOM 1063		ARG ARG	240	42.657	47.711	6.622	1.00	65.55
	ATOM 1064 ATOM 1065		ARG	240	41,828	48.040	7.073	1.00	0.00
50	ATOM 1065 ATOM 1066		ARG	240	43.298	47.122	7.115	1.00	0.00
	ATOM 1067	, M.22	ARG	240	38.863	49.791	2.164	1.00	41.91
	ATOM 1068	3 0	ARG	240	40,040	49.875	1.864	1.00	39.72
	ATOM 1069		GLU	241	37.969	49.138	1.389	1.00	46.63
55			GLU	241	37.022	49.233	1.606	1.00	0.00
٥.	ATOM 107		GLU	241	38.318	48.416	0.151	1.00	49.66
	ATOM 107		GLU	241	37.185	47.460	-0.280	1.00	58.65 76.02
	ATOM 107		GLU	241	37.559	46.447	-1.379	1.00	84.57
	ATOM 107		GLÜ	241	36.387	45.686	-2.058	1.00	86.53
6			GLU	241	35.539	46.310	-2.712 -1.970	1.00	90.34
-	ATOM 107		GLU	241	36.350	44.449	-1.009	1.00	45.34
	ATOM 107		GLU	241	38.559	49.403	-1.985	1.00	45.76
	ATOM 107		GLU	241	39.236	49.10Z 50.628	-0.966	1.00	40.75
	ATOM 107	19 N	GLY	242	38.015	20.020	0.550		

- 59 -

	ATOM	1080	н	GLY	242	37.428	50.949	-0.241	1.00	0.00
	ATOM	1081	CA	GLY	242	38.362	51.556	-2.014	1.00	35.23
	ATOM	1082	C	GLY	242	37.375	52.679	-2.047	1.00	33.55
5	ATOM	1083	0	GLY	242	36.694	52.955	-1.071	1.00	32.49
	ATOM	1084	N	ILE	243	37.290	53.311	-3.213	1:00	31.51
	ATOM ATOM	1085	H	ILE	243	37.707	52.888	-3.992	1.00	0.00
		1086	CA	ILE	243	36.590	54.576	-3.407	1.00	30.20
	ATOM	1087	CB	ILE	243	37.364	55.376	-4.511	1.00	31.68
10		1088	CG2	ILE	243	36.739	56.740	-4.856	1.00	28.31
10	ATOM	1089 1090	CĜI	ILE	243	38.749	55.674	-3.963	1.00	31.33
		1090	CDI	ILE	243	39.643	56.077	-5.135	1.00	35.46
		1092	C 0	ILE	243	35.148	54.304	-3.806	1.00	30.26
		1093	N	ILE	243	34.920	53.418	-4.618	1.00	30.91
15		1094	Н	CYS	244	34.151	55.017	-3.276	1.00	29.95
		1095	CA	CYS CYS	244	34.337	55.767	-2.666	1.00	0.00
		1096	CB		244	32.774	54.760	-3.652	1.00	31.93
	ATOM		SG	CYS CYS	244	31,855	55.223	-2.553	1.00	29.44
	ATOM		C	CYS	244	32.093	54.318	-1.030	1.00	36.45
20	ATOM		ō	CYS	244	32.291	55.415	-4.945	1.00	35.60
•	ATOM		N	GLY	244	32.427	56.619	-5.169	1.00	34.97
	ATOM		H	GLY	245 245	31.695	54.653	-5.863	1.00	34.56
	ATOM		CA	GLY	245 245	31.752	53.680	-5.763	1.00	0.00
	ATOM		C	GLY	245	30.973	55.279	-6. 9 62	1.00	37.28
25	ATOM	1104	ŏ	GLY	245	29.574	55.745	-6.552	1.00	42.13
	ATOM		N	LYS	246	29.085	55.433	-5.470	1.00	41.53
	ATOM		H	LYS	246	28.868 29.231	56.501	-7.415	1.00	44.81
	ATOM	1107	CA	LYS	246	27,604	56.669	-8.312	1.00	0.00
	ATOM	1108	CB	LYS	246	27.040	57.140 57.989	-7.033	1.00	47.47
30	ATOM	1109	CG	LYS	246	26.636	57.098	-8.186 -9.348	1.00	46.88
	ATOM		CD	LYS	246	25.882	57.806	-9.348 -10.459	1.00	51.88
	ATOM	1111	CE	LYS	246	24.560	58.422	-10.000	1.00 1.00	62.88
	MOTA		NZ	LYS	246	24.007	59.246	-11.063	1.00	69.93
	ATOM		HZI	LYS	246	23.833	58.663	-11.905	1.00	76.93 0.00
35	ATOM		HZ2	LYS	246	24.678	60.006	-11.295	1.00	0.00
	ATOM		HZ3	LYS	246	23,112	59.670	-10.742	1.00	0.00
	ATOM		С	LYS	246	26.477	56.209	-6.574	1.00	48.40
		1117	0	LYS	246	25.597	56.538	-5.7 78	1.00	42,25
40	ATOM		N	LYS	247	26.479.	54.982	-7.079	1.00	45.29
40	ATOM		H	LYS	247	27.176	54.727	-7.713	1.00	0.00
		1120	CA	LYS	247	25.465	54.045	-6.685	1.00	49,00
		1121	CB	LYS	247	25.209	53.130	-7.875	1.00	54.11
		1122	CG	LYS	247	24.508	53.886	-9.004	1.00	62,74
45		1123	CD	LYS	247	24.243	53.001	-10.220	1.00	72.81
43		1124	CE	LYS	247	23.531	53.721	-11.379	1.00	81.38
	MOTA	1125	NZ	LYS	247	23.373	52.835	-12.526	1.00	86,55
			HZI	LYS	247	22.808	52.005	-12.261	1.00	0.00
	ATOM	1127	HZ2	LYS	247	24.310	52.525	-12.854	00.1	0.00
50		1129	HZ3	LYS	247	22.899	53.347	-13.297	1.00	0.00
50	ATOM		Ċ	LYS	247	25.856	53.257	-5.450	1.00	49.50
	ATOM		0	LYS	247	25.319	52.203	-5.182	1.00	51,73
	MOTA		N	HIS	248	26.793	53.66 1	-4.603	1.00	50.46
	ATOM		H	HIS	248	27.235	54.526	-4.728	1.00	0.00
55	ATOM		CA CB	HIS	248	27.155	52.799	-3.498	1.00	45.73
	ATOM		CG	HIS HIS	248	28.438	53.305	-2.813	1.00 '	40.60
	ATOM		CD2	HLS	248	28.924	52.330	-1.737	1.00	36.78
	ATOM		ND1	HIS	248	29.411	51.071	-1.973	1.00	33.70
	ATOM		HDI	HIS	248	28.983	52.505	-0.416	1.00	38.30
60	ATOM		CEL	HIS	248	28.728	53.315	0.070	1.00	0.00
	ATOM		NEZ	HIS	248	29.467	51.412	0.129	1.00	34.58
	ATOM		HE2	HIS	248	29.720	50.547	-0.817	1.00	33.39
	ATOM		C	HIS	248	30.051	49.640	-0.658	1.00	0.00
	ATOM		ō	HLS	248	26.055	52.703	-2.473	1.00	48.32
			•	nω	248	25.591	53.685	-1.918	1.00	44.66

			650	249	25.626	51.478	-2.198	1.00	53.23
	ATOM 1144	N	SER SER	249	25.852	50.711	-2.770	1.00	0.00
	ATOM 1145	H	SER	249	24.798	51.233	-1.040	1.00	57.14
	ATOM 1146	CA CB	SER	249	23.445	50.596	-1.432	1.00	\$6.80
_	ATOM 1147	OG	SER	249	23.488	49.267	-1.938	1.00	58.69
5	ATOM 1148 ATOM 1149	HG	SER	249	23.996	49.345	-2.757	1.00	0.00
	ATOM 1149 ATOM 1150	C	SER	249	25.626	50.270	-0.262	1.00	59.00
	ATOM 1151	ō	SER	249	26.610	49.741	-0.760	1.00	55.94 68.05
	ATOM 1152	N	GLU	250	25.285	50.003	0.982	1.00	0.00
10	ATOM 1153	н	GLU	250	24.519	50.460	1.391	1.00	77.66
10	ATOM 1154	CA	GLU	250	25.982	48.958	1.710	1.00 1.00	85.81
	ATOM 1155	CB	GLU	250	25.518	48.929	3.169	1.00	96.97
	ATOM 1156	CG	GLU	250	25.746	50.253	3.922	1.00	103.17
	ATOM 1157	CD	GLU	250	25.505	50.056	5.411 6.085	1.00	108.23
15	ATOM 1158	OEL	GLU	250	25.196	51.045	5.895	1.00	105.30
	ATOM 1159	OE2	GLU	250	25.633	48.922	1.115	1.00	77.89
	ATOM 1160	С	GLU	250	25.778	47.573 46.720	1.119	1.00	78.08
	ATOM 1161	0	GLU	250	26.649	45.120 47.369	0.595	1.00	77.62
	ATOM 1162	N	GLN	251	24.574	48.135	0.546	1.00	0.00
20	ATOM 1163	H	GLN	251	23.971 24.155	46.078	0.085	1.00	77.63
•	ATOM 1164	CA	GLN	251		45.902	0.224	1.00	86.21
	ATOM 1165	СВ	GLN	251	22.616 21.677	47.096	0.015	1.00	97.44
	ATOM 1166	CG	GLN	251	21.754	48.076	1.182	1.00	106.14
	ATOM 1167	CD	GLN	251	22.556	49.005	1.219	1.00	112.85
25	ATOM 1168	OEI	GLN	251 251	20.940	47.965	2.214	1.00	108.86
	ATOM 1169	NE2	GLN	251	20.288	47.236	2.234	1.00	0.00
	ATOM 1170	HE21	GLN GLN	251	21.031	48.635	2.926	1.00	0.00
	ATOM 1171	HE22	GLN	251	24,544	45.830	-1.356	1,00	74.59
	ATOM 1172	С О.	GLN	251	24.771	44,704	-1 <i>.7</i> 78	1.00	73.25
30	ATOM 1173	N	VAL	252	24.636	46.863	-2,185	1.00	70.06
	ATOM 1174 ATOM 1175	H	VAL	252	24.241	47.718	-1.911	1.00	0,00 66,24
	ATOM 1175 ATOM 1176	CA	VAL	252	25.375	46.711	-3.425	1.00	68.01
	ATOM 1177	CB	VAL	252	24.482	46.902	-4.657	1.00	67.58
35	ATOM 1178	CG1	VAL	252	25.271	47.177	-5.941	1.00	71.77
دد	ATOM 1179	CG2	VAL	252	23.711	45.600	-4.832	1.00 1.00	60.42
	ATOM 1180	С	VAL	252	26.466	47.745	-3.424 -3.40 9	1.00	57.54
	ATOM 1181	0	VAL	252	26.282	48.961	-3.427	1.00	57.04
	ATOM 1182	N	PRO	253	27.648	47.275	-3.278	1.00	57.51
40		CD	PRO	253	28.022	45.875	-3.455	1.00	53.11
	ATOM 1184	CA	PRO	253	28.812	48.113 47.251	-2.798	1.00	57.01
	ATOM 1185		PRO	253	29,905	46.035	-2.272	1.00	57.83
	ATOM 1186		PRO	253	29.165	48.551	-4.866	1.00	47.52
	ATOM 1187		PRO	253	29.113 28.988	47,837	-5.853	1.00	45.85
45			PRO	253	29.533	49.808	-4.931	1.00	42.76
	ATOM 1189		ASP	254	29.509	50.376	-4.143	1.00	0.00
	ATOM 1190		ASP	254 254	30.038	50.374	-6.156	1.00	38.48
	ATOM 119		ASP	254	29.051	51.461	-6.580	1.00	36.32
_	ATOM 119		ASP	. 254	29.341	51.949	-7.979	1.00	36.27
5(O ATOM 119		asp Asp	254	28.990	53.090	-8.264	1.00	39.87
	ATOM 119		ASP	254	29.903	51.204	-8.786	1.00	36.65
	ATOM 119	5 OD2	ASP	254	31.400	50.909	-5.739	1.00	35.50
	ATOM 119	6 C 7 D	ASP	254	31.506	52.050	-5.321	1.00	34.98
-	ATOM 119	8 N	ILE	255	32.448	50.092	-5.R 37	1.00	33.54
5	5 ATOM, 119	9 H	ILE	255	32.350		-6.329	1.00	0.00
	ATOM 119	19 II 10 CA	ILE	255			-5.307	1.00	35.65
	ATOM 120	n CA	ILE	255			-4.295	1.00	39.66
	ATOM 120 ATOM 120		ILE	255			-3.797	1.00	40.81
_			ILE	255		49.254	-3.144	1.00	39.28 40.52
6			ILE	255	32.955			1.00	40.32 39.9 9
	• • • • • • • • • • • • • • • • • • • •	-	ILE	255				1.00	40.26
	ATOM 12 ATOM 12	-	ILE	255				1.00 1.00	
	ATOM 12		LEU	256		2 51.480	-6.503	1.00	33.07
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. 61 -

	MOTA	1208	H	LEU	256	35.471	52.284	-5.964	1.00	0.00
	ATOM	1209	CA	LEU	256	168.88	51.410	-7.340	1.00	39.66
	ATOM	1210	CB	LEU	256	37.066	52.755	-8.074	1.00	38.42
_	ATOM	1211	CG	LEU	256	38.298	52.842	-8.988	1.00	37.25
5		1212	CDI	LEU	256	38.214	51.825	-10.115	1.00	36.68
		1213	CD2	LEU	256	38.377	54.245	-9.571	1.00	39.39
	ATOM	1214	C	LEU	256	38.022	51.095	-6.407	1.00	38.52
		1215	0	LEU	256	38.368	51.793	-5.455	1.00	40.24
10		1216	N	GLN	257	38.658	49.977	-6.715	1.00	39.82
10	MOTA MOTA	1217	H	GLN	257	38.308	49.468	-7.488	1.00	0.00
		1219	CB CA	GLN	257	39.795	49.436	-5.992	1.00	42.81
	ATOM		CG	GLN GLN	257	40.050	48.020	-6.455	1.00	49.86
		1221	CD	GLN	257	40.720	47.077	-5.487	1.00	62.03
15		1222	OEI	GLN	257 257	39.707	46.054	-5.046	1.00	66.77
		1223	NE2	GLN	257	38.568 40.013	46.361	-4.726	1.00	70.03
		1224	HE21	GLN	257	40.813	44.767 44.483	-5.063 5.260	1.00	67.84
		1225	HE22	GLN	257	39.295	44.174	-5.369 -4.752	1.00	0.00
		1226	C	GLN	257	40.991	50.253	-6.300	1.00 1.00	9.00 43.83
20		1227	0	GLN	257	41.296	50.526	-7.457	1.00	44.09
•	ATOM	1228	N	LEU	258	41.747	50.678	-5.315	1.00	45.79
		1229	H	LEU	258	41.461	50.433	-4.414	1.00	0.00
		1230	CA	LEU	258	42.951	51.468	-5.556	1.00	48.19
	ATOM		CB	LEU	258	43.638	51.714	-4.187	1.00	62.23
25	ATOM		CG	LEU	258	45.178	51.726	-4.015	1.00	71.72
	ATOM		CD1	LEU	258	45.742	53.135	-3.795	1.00	65.86
	ATOM		CD2	LEU	258	45.495	50.830	-2.809	1.00	80.83
	MOTA		C	LEU	258	43.958	50.864	-6.564	1.00	46.26
30		1236	0	LEU	258	44.583	51.564	-7.368	1.00	43.62
30		1237 1 23 8	N H	ASN	259	44.214	49.561	-6.654	1.00	41.70
		1239	CA	ASN ASN	259	43.687	48.856	-6.205	1.00	0.00
		1240	CB	ASN	259 259	45.197	49.203	-7.648	1.00	41.88
		1241	CG	ASN	259	46.029 45.376	48.071	-7.151	1.00	46.19
35	ATOM	1242	OD1	ASN	259	44.187	46.750 46.674	-7.280 -7.008	1.00 1.00	49.87
		1243	ND2	ASN	259	46.115	45.710	-7.644	1.00	56.41 50.33
	MOTA	1244	HD21	ASN	259	47.075	45.857	-7.810	1.00	0.00
	MOTA	1245	HD22	ASN	259	45.682	44.830	-7.714	1.00	0.00
	MOTA	1246	С	ASN	259	44.636	48.900	-9.007	1.00	38.03
40		1247	0	ASN	259	45.417	48.585	-9.896	1.00	35.47
		1248	N	ALA	260	43.309	48.952	-9.180	1.00	33.64
	ATOM	1249	H	ALA	260	42.708	49.000	-8.409	1.00	0.00
		1250	CA	ALA	260	42.732	48.991	-10.512	1.00	29.46
45		1251	CB	ALA	260	41.224	49.011	-10.356	1.00	25.72
45	MOTA MOTA	1252	C	ALA	260	43.243	50.221	-11.230	1.00	28.23
		1253 1254	O N	ALA	260	43.637	50.176	-12.385	1.00	31.28
	ATOM	1255	H	ile Ile	261	43.255	51.359	-10.532	1.00	28.30
	ATOM	1256	CA	LE	261 251	42.856 43.812	51.344 52.619	-9,640	1.00	0.00
50	ATOM	1257	СВ	ILE	· 261	43.758	53.698	-11.027 -9.884	1.00 1.00	25.68
		1258	CG2	ILE	261	44.364	55.014	-10.371	1.00	27.06 22.81
	MOTA		CGI	ILE	261	42.300	53.941	-9.444	1.00	23.10
	MOTA		CDI	ILE	261	42.175	54.847	-8.215	1.00	15.19
	MOTA	1261	С	ILE	261	45.246	52.443	-11.516	1.00	27.03
55	MOTA	1262	0	ILE.	261	45.594	52.815	-12.626	1.00	30.14
	MOTA	1263	N	PHE	262	46.130	51.861	-10.699	1.00	25.11
		1264	H	PHE	262	45.832	51.647	-9.791	1.00	0.00
	MOTA		CA	PHE	262	47.511	51.582	-11.075	1.00	23.29
		1266	CB	PHE	262	48.275	50.853	-9.941	1.00	23.94
60	ATOM	1267	CG	PHE	262	48.795	51.855	-8.914	1.00	22.84
		1268	CDI	PHE	262	47.919	52.571	-8.110	1.00	26.50
		1269	CD2	PHE	262	50.147	52.112	-8.831	1.00	22.06
	ATOM	1270	CEI	PHE	262	48.402	53.570	-7.279	1.00	29.99
	ATOM	1271	CE2	PHE	262	50.619	53.093	-7.974	1.00	24.85

									00.46
	. mov. / 1373	cz	PHE	262	49.756	53.842	-7.209	1.00	20.46
	ATOM 1272	C	PHE	262	47.573	50.725	-12.288	1.00	27.29
	ATOM 1273	Ö	PHE	262	48.285	50.994	-13.247	1.00	29.20
	ATOM 1274		ASN	263	46.796	49.640	-12.270	1.00	29.26
_	ATOM 1275	N	ASN	263	46.219	49.470	-11.496	1.00	0.00
5	ATOM 1276	H			46.827	48.683	-13.372	1.00	32.07
	ATOM 1277	CA	ASN	263	45.872	47.495	-13.119	1.00	36.54
	ATOM 1278	CB	ASN	263	46.483	46.467	-12.141	1.00	48.06
	ATOM 1279	CG	ASN	263	47.677	46.416	-11.842	1.00	52.97
	ATOM 1280	ODI	ASN	263	47.677	45.600	-11.517	1.00	50.97
10	ATOM 1281	ND2	ASN	263		45.639	-11.681	1.00	0.00
	ATOM 1282	HD21	ASN	263	44.728	44.947	-10.928	1.00	0.00
	ATOM 1283	HD22	ASN	263	46.129	49.332	-14.673	1.00	31.78
	ATOM 1284	С	ASN	263	46.450	49.081	-15.715	1.00	30.44
	ATOM 1285	0	ASN	263	47.043	50.203	-14.640	1.00	30.71
15	ATOM 1286	N	MET	264	45.443		-13.807	1.00	0.00
	ATOM 1287	H	MET	264	44.940	50.348	-15.817	1.00	31.45
	ATOM 1288	CA	MET	264	45.054	50.956	-15.565	1.00	36.38
	ATOM 1289	CB	MET	264	43.742	51.698	-15.835	1.00	44.27
	ATOM 1290	CG	MET	264	42.552	50.779	-15.768	1.00	51.17
20	ATOM 1291	SD	MET	264	40.953	51.627	-14.191	1.00	51.48
	ATOM 1292	CE	MET	264	40.447	51.015		1.00	30.63
	ATOM 1293	C	MET	264	46.036	51.979	-16.362	1.00	25.16
	ATOM 1294	Ó	MET	264	45,903	52.396	.17.500	1.00	27.60
	ATOM 1295	N	LEU	265	47.044	52.435	-15.617	1.00	0.00
25	ATOM 1296	H	LEU	265	47.167	52.116	-14.695	1.00	23.27
23	ATOM 1297	CA	LEU	265	47.941	53.456	-16.133	1.00	24.95
	ATOM 1298	CB	LEU	265	47.844	54.709	-15.272	1.00	30.77
	ATOM 1299		LEU	265	46.508	55.432	-15.452	1.00	35.13
	ATOM 1300		LEU	265	45.942	55.833	-14.099	1.00	30.86
.30	ATOM 1301		LEU	265	46.722	56.641	-16.339	1.00	21.55
٠,٥٠	ATOM 1302		LEU	265	49.365	52.980	-16.157		21.21
	ATOM 1303		LEU	265	50.309	53.724	-16.327	1.00	20.49
	ATOM 130		NZA	266	49.583	51. 69 6	-15.972	1.00	0.00
	ATOM 130		ASN	266	48.832	51.117	-15.737	1.00	21.68
35			ASN	266	50.918	51.154	-16.072	1.00	20.66
35			ASN	266	50.905	49.810	-15.355	1.00	26.15
	ATOM 130 ATOM 130		ASN	266	50.208	48. 69 2	-16.103	1.00	30.13
			ASN	266	50.004	48.725	-17.302	1.00	26.18
			ASN	266	49.889	47.599	-15.435	1.00	0.00
40			ASN	266	50.148	47.527	-14.492	1.00	0.00
40			ASN	266	49.389	46.885	-15.880	1.00	27,52
			ASN	266	51.463	51.028	-17.496	1.00	
	ATOM 131		ASN	266	50.834	51.391	-18.480	1.00	25.25
	ATOM 131		THR	267	52.671	50.496	-17. 66 8	1.00	29.62
	ATOM 131		THR	267	53.155	50.095	-16.911	1.00	0.00
4:	5 ATOM 13	10 A	THR	267	53.325	50.491	-18.965	1.00	31.01
	ATOM 13		THR	267	54.807	50.073	-18.787	1,00	31.03
	ATOM 13		THR	267	55.330	51.030	-17.858	1.00	36.10
	ATOM 13			267	54.960	50.998	-16.964	1.00	0.00
	ATOM 13		THR	. 267	55.686	50.139	-20.052	1.00	28.06
5	0 ATOM 13		THR	•	52.618	49.591	-19.933	1.00	31.70
	ATOM 13	22 C	THR	267	52.606	49,803	-21.128	1.00	31.87
	ATOM 13	23 O	THR	267	51.988	48.543	-19.427	1.00	34.40
	ATOM 13	124 N	LYS	268	52.010		-18.464	1.00	0.00
	ATOM 13	325 H	LYS	268			-20.275	1.00	34.41
5	5 ATOM I	326 CA	LYS	268	51.228		-19.458	1.00	38.03
	ATOM 13		LYS	268				1.00	46.78
	ATOM I	328 CG	LYS	268				1.00	51.28
	ATOM I		LYS	268				1.00	59.23
	ATOM I		LYS	268				1.00	63.39
_	O ATOM 1		LYS	268				1.00	0.00
•	ATOM 1		LYS	268				1.00	0.00
	ATOM 1			268					0.00
	ATOM 1			268	47.263				34.09
	ATOM I	1335 C	LYS	26		3 48.367	7 -20.803	1.00	J4.U
	Alum	ک روز		34					

	ATOM		0	LYS	268	49.735	48.325	-21.979	1.00	34.97
	ATOM		N	ASN	269	49.267	49.054	-19.948	1.00	29.59
	MOTA	1338	H	A5N	269	49.577	49,149	-19.020	1.00	0.00
5	MOTA		CA	ASN	269	48.012	49.651	-20.366	1.00	24.54
3	MOTA MOTA		CB	ASN	269	46.995	49.607	-19.263	1.00	26.50
		1341	CG	ASN	269	46.685	48.179	-19.067	1.00	32.19
		1343	ODI	ASN	269	46.295	47.488	-19.988	1.00	35.32
	ATOM		ND2 HD21	ASN ASN	269	46.896	47.620	-17.892	1.00	35.76
10		1345	HD22	asn Asn	269	47.263	48.175	-17.168	1.00	0.00
	ATOM		C	ASN	269 269	46.674 48.130	46.672	-17.821	1.00	0.00
	ATOM	1347	o.	ASN	269	47.253	51.078 51.677	-20.767	1.00	27.02
		1348	N	CYS	270	49.228	51.744	-21.370 -20.458	1.00	27.85
	MOTA	1349	H	CYS	270	49.948	51.324	-19.931	1.00 1.00	26.07 0.00
15	MOTA	1350	CA	CYS	270	49.351	53.133	-20.848	1.00	23.70
		1351	CB	CYS	270	49.028	54.048	-19.676	1.00	22.29
		1352	SG	CYS	270	48.971	55.800	-20.133	1.00	27.75
		1353	C	CYS	270	50.770	53.374	-21.287	1.00	26.34
20		1354	0	CYS	270	51.515	54.159	-20.711	1.00	24.34
2,0	MOTA		N	PRO	271	51.237	52.761	-22.313	1.00	26.04
		1356	CD	PRO	271	50.428	52.006	-23.263	1.00	26.80
		1357 1358	CA	PRO	271	52.628	52.860	-22.730	1.00	26.32
		1359	CB CG	PRO	271	52.694	51.997	-23.974	1.00	25.30
25		1360	C	PRO PRO	271	51.292	52.036	-24.514	1.00	24.28
		1361	ò	PRO	271	53.104	54,281	-22.955	1.00	29.26
		1362	ĸ	SER	271 272	54.259	54.644	-22.776	1.00	31.29
		1363	H	SER	272	52.220 51.287	55.180 54.942	-23.365	1.00	30.94
	ATOM		CA	SER	272	52.617	56.555	-23.574 -23.545	1.00	0.00
30	ATOM	1365	CB	SER	272	51.440	57.305	-23.343 -24.135	1.00 1.00	31.44 37.05
	ATOM	1366	OG	SER	272	50.225	56.635	-23.790	1.00	53.17
	MOTA	1367	HG	SER	272	50.053	56.659	-22.840	1.00	0.00
	ATOM		С	SER	272	53.068	57.208	-22.269	1.00	30.10
2-	ATOM		0	SER	272	53.666	58.259	-22.345	1.00	30.70
35	MOTA		N	LEU	273	52.826	56.687	-21.064	1.00	29.16
	ATOM		H	LEU	273	52.343	55.839	-20.921	1.00	0.00
	MOTA MOTA		CA CB	LEU LEU	273	53.350	57.384	-19.910	1.00	26.90
		1374	CG	LEU	273	52.250	57.567	-18.851	1.00	23.25
40	ATOM		CD1	LEU	273 273	51.343	58.788	-19.139	1.00	24.56
	MOTA		CD2	LEU	273	50.112 52.168	58.798 60.056	-18.225 -18.965	1.00	20.80
	ATOM		C	LEU	273	54.519	56.632	-19.336	1.00 1.00	20.75 28.09
	ATOM	1378	0	LEU	273	54.948	56.803	-18.205	1.00	24.97
	ATOM	1379	N	LYS	274	55.108	55.731	-20.112	1.00	28.29
45		1380	H	LYS	274	54.753	55.544	-21.008	1.00	0.00
		1381	CA	LYS	274	56.314	55.053	-19.684	1.00	27.08
	ATOM		CB	LYS	274	56.853	54.212	-20.819	1.00	28.08
	MOTA		CG	LYS	274	58.111	53.417	-20.485	1.00	32.44
50	ATOM		CD	LYS	274	58.619	52.783	-21.777	1.00	40.58
50	MOTA MOTA		CE	LYS	274	59.960	52.073	-21.658	1.00	45.30
	ATOM		NZ HZ1	LYS LYS	274	61.040	53.035	-21.669	1.00	47.25
	ATOM		HZ2	LYS	274 274	60.925	53.685	-20.867	1.00	0.00
	ATOM		HZ3	LYS	274	61.021 61.949	53.571 53.571	-22.560	1.00	0.00
55	ATOM		C	LYS	274	57.348	52.537 56.087	•21.580 •19.285	1.00 1.00	0.00
	ATOM		ō	LYS	274	57.594	57.057	-19.983	1.00	27.81 27.52
	ATOM		N	ASP	275	57.942	55.858	-19.303	1.00	27.52 26.78
	MOTA	1393	H	ASP	275	57.514	55.152	-17.599	1.00	0.00
_	ATOM		CA	ASP	275	59.007	56.689	-17.546	1.00	24.40
60	MOTA		CB	ASP	275	60.222	56.827	-18.507	1.00	24.70
	ATOM		CG	ASP	275	60.849	55.482	-18.831	1.00	28.62
	MOTA		ODI	ASP	275	61.282	55.358	-19.970	1.00	35.31
	MOTA		OD2	ASP	275	60.919	54.577	-17.980	1.00	31.34
	MOTA	1399	С	ASP	275	58.618	58.090	-17.114	1.00	21.07

	ATOM 1400	0	ASP	275	59.450	.58.931	-16.787	1.00	22.30
	ATOM 1401	N	LYS	276	57.332	58.409	-17.087	1.00	20.11
	ATOM 1402	Н	LYS	276	56.635	57.754	-17.318	1.00	0.00
	ATOM 1403	CA	LYS	276	56.918	••••	-16.690	1.00	20.03
5	ATOM 1404	СВ	LYS	276	55.977	60.263	-17.772	1.00	15.62 20.79
_	ATOM 1405	CG	LYS	276	56.700	60.352	-19.129	1.00 1.00	25.10
	ATOM 1406	CD	LYS	276	55.762	60.741	-20.275 -21.682	1.00	24.43
	ATOM 1407	CE	LYS	276	56.341	60.584	-21.682 -22.677	1.00	26.93
	ATOM 1408	NZ	LYS	276	55.317	60.858 61.828	-22,562	1.00	0.00
10	ATOM 1409	HZ1	LYS	276	54.962 54.528	60.192	-22.550	1.00	0.00
	ATOM 1410	HZ2	LYS LYS	276 276	55.716	60.744	-23.630	1.00	0.00
	ATOM 1411	HZ3	LYS	276	56.238	59.664	-15.339	1.00	21.33
	ATOM 1412 ATOM 1413	С О	LYS	276	55.549	58.696	-15.080	1.00	23.74
15	ATOM 1414	N	PRO	277	56.342	60.565	-14.432	1.00	18.25
13	ATOM 1415	CD	PRO	277	57.125	61.779	-14.555	1.00	20.11
	ATOM 1416	CA	PRO	277	55.658	60.491	-13.158	1.00	19.61
	ATOM 1417	CB	PRO	277	56.178	61.665	-12.348	1.00	19.47 17.30
	ATOM 1418	CG	PRO	277	57.365	62.160	-13.113	1.00 1.00	21.46
20	ATOM 1419	С	PRO	277	54.146	60.506	-13.252 -13.988	1.00	24.07
•	ATOM 1420	0	PRO	277	53.522	61.268	-12,473	1.00	19.22
	ATOM 1421	N	LYS	278	53,531	59.642 58.986	-11.955	1.00	0.00
	ATOM 1422	H	LYS	278 278	54.049 52.093	59.598	-12.436	1.00	18.47
	ATOM 1423	CA	LYS LYS	278	51.654	58.233	-12.953	1.00	16.94
25	ATOM 1424	CB CG	LYS	278	52.074	58.090	-14.435	1.00	15.20
	ATOM 1425 ATOM 1426	CD	LYS	278	51.919	56.673	-14. 9 45	1.00	16.03
	ATOM 1427	CE	LYS	278	53.057	55.723	-14.550	1.00	18.36
	ATOM 1428	NZ	LYS	278	54.258	55.975	-15.318	1.00	19.84 0.00
30	ATOM 1429	HZI	LYS	278	54.573	56.952	-15.158	1.00 1.00	0.00
• :	ATOM 1430	HZ2	LYS	278	54.052	55.841	-16.328	1.00	0.00
	ATOM 1431	HZ3	LYS	278	55.008	55.317	-1 5.025 -11.016	1.00	21.76
	ATOM 1432	C	LYS	278	51.698	59.807 58.916	-10.202	1.00	23.16
	ATOM 1433	0	LYS	278	51.903 51.133	60.956	-10.655	1.00	18.92
35	ATOM 1434	N	VAL	279	50.893	61.619	-11.338	1.00	0.00
	ATOM 1435	H	VAL VAL	279 279	50.830	61.249	-9.270	1.00	16.68
	ATOM 1436	CA CB	VAL	279	51.262	62.707	-8.960	1.00	15.16
	ATOM 1437 ATOM 1438	CGI	VAL	279	51.042	63.041	-7.484	1.00	12.89
40		CG2	VAL	279	52.737	62.882	-9.318	1.00	13.20 20.13
40	ATOM 1440	C	VAL	279	49.340	61.054	-9.066	1.00	20.13
	ATOM 1441	0	VAL	279	48.520	61.647	-9.770 P. 000	1.00 1.00	20.07
	ATOM 1442	N	ILE	280	48.946	60.212	-8.099 -7.543	1.00	0.00
	ATOM 1443		ILE	280	49.628	59.772 59.939	-7.830	1.00	20.53
45			ILE	280	47.535 47.306	58.401	-7.914	1.00	25,21
	ATOM 1445		ILE	280	45.900	58.000	-7,454	1.00	17.77
	ATOM 1446		ILE	280 280	47.519	57.977	-9.377	1.00	24.13
	ATOM 1447		ILE ILE	280	48.045	56.562	-9,513	1.00	30.13
- /	ATOM 1448		ILE	280	47.150	60,496	-6.467	1.00	21.62
5.0	ATOM 1449 ATOM 1450		ILE	280	47.789	60.204	-5.464	1.00	22.51
	ATOM 1450	N	ILE	281	46.108	61.318	-6,363	1.00	20.14
	ATOM 145	2 H	ILE	281	45.542	61.495	-7.148	1.00	0.00
	ATOM 145		ILE	281	45.770	61.940	-5.109	1.00	19.43 19.82
5			ILE	281	45.858	63.465	-5.300	1.00 1.00	16.34
	ATOM 145		ILE	281	45.338	64.229	-4.081 5.516	1.00	24.03
	ATOM 145		ILE	281	47,333	63.810	-5.516 -5.781	1.00	26.46
	ATOM 145	7 CD1	ILE.	281	47.627	65.269	-3.781 -4.737	1.00	22.29
	ATOM 145		ILE	281	44.389		-5.568	1.00	21.67
6	O ATOM 145	9 0	ILE	281	43.495		-3.509	1.00	23.39
	ATOM 146		ILE	282	44.128 44.833			1.00	0.00
	ATOM 146		ILE	282	42.781			1.00	24.32
	ATOM 140		ILE	282				1.00	21.45
	ATOM 144	63 CB	ILE	282	42.701	57.454			

	ATOM	1464	CG2	ILE	282	41.368	58.517	-2.706	1.00	22.39
	ATOM	1465	CG1	ILE	282	43,224	58.492	-4.387	1.00	17.29
	ATOM	1466	CDI	ILE	282	43.645	57.044	-4.387	1.00	18.62
	MOTA	1467	С	ILE	282	42.275	61.214	-1.878	1.00	24.17
5	ATOM	1468	0	ILE	282	42.837	61.022	-0.802	1.00	23.02
	ATOM	1469	N	GLN	283	41.195	61.985	-1.981	1.00	24.17
	ATOM	1470	H	GLN	283	40.880	62.229	-2.879	1.00	0.00
	ATOM		CA	GLN	283	40.417	62.361	-0.807	1.00	26.94
	ATOM	1472	CB	GLN	283	39.916	63.809	-0.951	1.00	
10	ATOM		CG	GLN	283	38.790	64.295	-0.931	1.00	29.78
		1474	CD	GLN	283	39.213	64.338	1.454	1.00	30.76
	ATOM		OÉI	GLN	283	40.263	64.820	1.835		31.96
		1476	NEZ	GLN	283	38.367			1.00	26.84
	ATOM		HE21	GLN	283	37.468	63.923	2.370 2.058	1.00	32.78
15		1478	HE22	GLN	283	38.632	63.628	3.299	1.00	0.00
	ATOM		C	GLN	283	39.227	63.911		1.00	0.00
	ATOM		ŏ	GLN	283		61.397	-0.673	1.00	27.75
	ATOM	1481	N	ALA	284	38.396	61.282	-1.580	1.00	27.45
	ATOM		H	ALA	284	39.127	60.692	0.460	1.00	24.57
20	ATOM	1483	CA	ALA		39.827	60.742	1.148	1.00	0.00
٠,٠	ATOM		CB	ALA	284	38.115	59.682	0.688	1.00	22.92
	ATOM	1485	C	ALA	284	38.196	58.595	-0.383	1.00	15.36
	ATOM		0		284	38.438	59.060	2.029	1.00	25.81
	ATOM		N	ALA ALD	284	39.598	58.871	2.387	1.00	29.97
25	ATOM		CA		285	37.396	58.754	2.795	1.00	27.43
2	ATOM			ALD	285	37.538	58.081	4.104	1.00	26.59
	MOTA		С О	ALD	285	38.038	56.689	3.860	1.00	26.61
	ATOM			ALD	285	37.779	56.155	2.790	1.00	30.02
		1492	CB SG	ALD ALD	285	36.215	57.908	4.846	1.00	26.43
30	ATOM				285	35.603	59.475	5.487	1.00	34.69
30	ATOM		NI	ALD	285	32.516	69.905	10.B51	1.00	58.66
	ATOM		Cl	ALD	285	33.306	68.911	10.160	1.00	48.23
	ATOM		· Cs	ALD	285	32.529	67.649	10.069	1.00	45.89
			O2	ALD	285	31.833	67.247	10.983	1.00	47.20
35	MOTA		C9	ALD	285	34.623	68.691	10.902	1.00	46.18
33	MOTA		CG	ALD	285	35.549	69.731	10.350	1.00	50.40
	MOTA		CD1	ALD	285	35.586	71.002	10.923	1.00	53.28
	MOTA		CD2	ALD	285	36.268	69.456	9.185	1.00	53.34
	MOTA		CEI	ALD	285	36,376	71.984	10.326	1.00	55.15
40	ATOM		CE2	ALD	285	37.056	70.446	8.595	1.00	52.67
40	ATOM		CZ	ALD	285	37.109	71.725	9.163	1.00	57.31
	ATOM		OH	ALD	285	37.756	72.792	8.546	1.00	63.58
	MOTA	1505	N2 ~	ALD	285	32.657	67.034	8.925	1.00	38.70
	MOTA		C2	ALD	285	31.932	65.809	8.702	1.00	34.11
45		1507	C6	ALD	285	33.033	64.915	8.206	1.00	33.25
45	ATOM	1508	03	ALD	285	34.021	65.383	7.657	1.00	27.84
	ATOM		C10	YTD	285	30.819	66.116	7.661	1.00	39.67
	ATOM		CG1	ALD	285	30.399	64.915	6.822	1.00	44.49
	ATOM		CG2	YLD	285	29.588	66.547	8.453	1.00	40.46
- 0	ATOM	1512	N3	ALD	285	32.851	63.630	8.422	1.00	31.01
50	ATOM	1513	C3	ALD	. 285	33.824	62.646	7.997	1.00	33,50
	ATOM	1514	C7	ALD	285	33.316	62.147	6.681	1.00	37.64
	ATOM		04	ALD	285	32.114	62.219	6.432	1.00	44.05
	MOTA		CII	ALD	285	33.899	61.464	8.952	1.00	31.45
	ATOM		N4	ALD	285	34.265	61.650	5.908	1.00	35.48
55	ATOM		C4	ALD	285	34.210	61.553	4.464	1.00	33.98
	MOTA		C8	ALD	285	34.763	60.203	4.008	1.00	36.52
	ATOM		C12	ALD	285	35.030	62.710	3.971	1.00	35.04
	MOTA		C13	ALD	285	34.835	62.936	2.510	1.00	37.05
	ATOM		ODI	ALD	285	35.766	63.453	1.898	1.00	37.30
60	ATOM		OD2	ALD	285	33.772	62.594	1.993	1.00	34.87
	ATOM		05	ALD	285	33.751	59.337	3.478	1.00	42.82
	ATOM		HI	ALD	28.5	36.497	58.977	2.418	1.00	0.00
	MOTA		C14	YTD	285	31.983	70.803	10.042	1.00	65.26
	ATOM	1527	H5	ALD	285	32.384	69.885	11.844	1.00	0.00

								4.00	0.00
	ATOM 1528	Н6	ALD	285	37.739	72.711	7.587	1.00 1.00	0.00
	ATOM 1529		ALD	285	33.249	67.364	8.179 8.842	1.00	0.00
	ATOM 1530	H8	ALD	285	32.004	63.310	6.347	1.00	0.00
	ATOM 1531	H9	ALD	285	35.098	61.322 70.799	10.002	1.00	65.03
5	ATOM 1532	C15	ALD	285	30.476 32.684	71.569	9.386	1.00	73.24
	ATOM 1533	01	ALD	285 285	35.481	60.411	3.215	1.00	0.00
	ATOM 1534	H2	ALD ARG	286	38.740	56.061	4.793	1.00	26.38
	ATOM 1535	N H	ARG	286	38.960	56.493	5.651	1.00	0.00
٦.	ATOM 1536 ATOM 1537	CA	ARG	286	39.231	54.719	4,550	1.00	25.85 24.02
10	ATOM 1538	CB	ARG	286	40.747	54.692	4.682	1.00 1.00	23.75
	ATOM 1539	CG	ARG	286	41.387	55.701	3.739 2.926	1.00	27.07
	ATOM 1540	CD	ARG	286	42.430	54.961	3.356	1.00	29.64
	ATOM 1541	NE	ARG	286	43.768	55.251 55.944	4.031	1.00	0.00
15	ATOM 1542	HE	ARG	286	43.926 44.798	54.588	2.845	1.00	26.29
	ATOM 1543	CZ	ARG	286 286	46.050	55.018	3.144	1.00	25.89
	ATOM 1544	NHI	ARG ARG	286	46.181	55.814	3.735	1.00	0.00
	ATOM 1545 ATOM 1546	HHII HHI2	ARG	286	46.845	54.538	2.774	1.00	0.00 19.53
20	ATOM 1547	NH2	ARG	286	44,621	53.491	2.058	1.00 1.00	0.00
2,0	ATOM 1548	HH21	ARG	286	43.700	53.161	1.851 1.689	1.00	0.00
	ATOM 1549	HH22	ARG	286	45.416	53.010	5.538	1.00	26.43
	ATOM 1550	С	ARG	286	38.629	53.755 52.683	5.803	1.00	27.12
	ATOM 1551	0	ARG	286	39.160 37.495	54.113	6.133	1.00	26.73
25	ATOM 1552	N	GLY	287 287	37.021	54.925	5.843	1.00	0.00
	ATOM 1553	H	GLY GLY	287	36.905	53.350	7.208	1.00	27.46
	ATOM 1554 ATOM 1555	CY CY	GLY	287	36.162	54.330	8.100	1.00	30.03 32.06
	ATOM 1555 ATOM 1556		GLY	287	35.932	35.475	7.728	1.00 1.00	33.55
30	ATOM 1557		ASP	288	35.804	53.848	9.292 9.545	1.00	0.00
-	ATOM 1558		ASP	288	36.263	53.019	10.273	1.00	32.84
	ATOM 1559		ASP	288	34.941	54.525 53.609	10.820	1.00	40.57
	ATOM 1560		ASP	288	33.839 33.043	52.985	9.706	1.00	48.86
	ATOM 1561		ASP ASP	288 288	33.051	51.752	9.612	1.00	53.99
35	ATOM 1562		ASP	288	32.413	53.731	8.944	1.00	51.26
	ATOM 1563 ATOM 1564		ASP	288	35.710	55.003	11.500	1.00	31.98 32.42
	ATOM 156		ASP	288	35.283	55.884	12.250	1.00 1.00	28.78
	ATOM 156		SER	239	36.884	54.437	11.775 11.200	1.00	0.00
40		7 H	SER	289	37.321	53.757 54.805	12.969	1.00	24.44
	ATOM 156		SER	289	37,596 38,704	53.801	13.071	1.00	22.45
	ATOM 156		SER	289 289	38.083	52.535	13.042	1.00	33.26
	ATOM 157		SER SER	289	38.773	51.861	13.003	1.00	0.00
4.5	ATOM 157		SER	289	38.069	56.239	12.940	1.00	24.99 26.10
45	5 ATOM 157 ATOM 157		SER	289	38.394	56.809	11.907	1.00 1.00	27.20
	ATOM 15	-	PRO	290	38.143	56.885	14.041	1.00	22.52
	ATOM 15		PRO	290	37.408	56.497	15.249 14.136	1.00	25.66
	ATOM 15		PRO	290	38.631	58.270 58.769	15.454	1.00	27.04
5	0 ATOM 15		PRO	290 290	38.051 37.920	57.494	16.266	1.00	21.06
	ATOM 15	78 CG	PRO	290 290	40.138	58.521	14.025	1.00	28.40
	ATOM 15	79 C	PRO PRO	290	40.637	59.638	13.939	1.00	26.84
	ATOM 15	80 O	GLY	291	40.904	57.441	14.034	1.00	24.70 0.00
_	ATOM 15		GLY	291	40.473	56.576	14.160	1.00	27.48
5	5 ATOM 15		GLY	291	42.342		13.835	1.00 1.00	26.97
	ATOM 1	584 C	GLY	291			15.056	1.00	24.56
	ATOM 1	585 O	GLY	291			14.877 16.287	1.00	24.36
	ATOM I		VAL	292			16.414	1.00	0.00
6	O ATOM 1	587 H	VAL	292			17.491	1.00	26.77
_	ATOM 1	588 CA	VAL	292 293			18.102		24.74
	ATOM I	.589 CB	VAL VAL	29.		44.	17.151	1.00	28.54
	ATOM I	1590 CG1 1591 CG		29:	_		18.371	1.00	25.92
	ATOM I	ואנו בענו	, ,,,,,						

	ATOM	1592	С	VAL	292	43.273	57.168	18.641	1.00	26.43
	ATOM	1593	0	VAL	292	42.441	56.276	18.749	1.00	28.53
	ATOM	1594	N	VAL	293	44.199	57.326	19.569	1.00	27.36
_	ATOM		H	VAL	293	44.833	58.079	19.525	1.00	0.00
5	ATOM		CA	VAL	293	44.287	56.466	20.719	1.00	28.84
	ATOM	1597	CB	VAL	293	45.426	55.456	20.369	1.00	27.36
	ATOM		CG1	VAL	293	46.801	55.914	20.801	1.00	23.55
	ATOM		CG2	VAL	293	45.069	54.152	21.025	1.00	28.81
10	ATOM		C	VAL	293	44.564	57.485	21.845	1.00	30.19
10	ATOM		0	VAL	293	45.149	58.538	21.595	1.00	30.10
	ATOM		N	TRP	294	44.153	57.245	23.098	1.00	27.38
	ATOM		H	TRP	294	43.657	56.424	23.290	1.00	0.00
	MOTA MOTA	1604 1605	CA	TRP	294	44.490	58.156	24.193	1.00	26.27
15	ATOM	1606	CB	TRP	294	43.553	58.075	25.389	1.00	24.27
	ATOM		CG CD2	TRP	294	42.160	58.474	25.030	1.00	26.12
	ATOM	1608	CE2	TRP TRP	294	41.796	59.897	24.914	1.00	27.34
	ATOM		CE3	TRP	294	40.345	59.680	24.556	1.00	25.56
	ATOM		CDI	TRP	294 294	42.296	61.171	25.019	1.00	28.09
20	ATOM	1611	NEI	TRP	294 294	41.128	57.603	24.792	1.00	25.19
•	ATOM		HE1	TRP	294	40.077	58.356	24.518	1.00	28.67
	MOTA	1613	CZ2	TRP	294	39.189 39.526	57.990	24.324	1.00	0.00
	ATOM	1614	CZ3	TRP	294	41.421	60.761	24.347	1.00	21.74
	MOTA	1615	CH2	TRP	294	40.084	62.218 62.017	24.795	1.00	27.80
25	MOTA	1616	С	TRP	294	45.823	57.790	24.478 24.766	1.00	25.43
	ATOM	1617	0	TRP	294	46.217	56.632	24.700	1.00 1.00	27.42
	ATOM	1618	N	PHE	295	46.574	58.751	25,251	1.00	26.36 32.18
	MOTA	1619	H	PHE	295	46.375	59.702	25.116	1.00	0.00
	MOTA	1620	CA	PHE	295	47.704	58.370	26.063	1.00	41.42
30	MOTA	1621	CB	PHE	295	49.007	58.377	25.207	1.00	40.27
	ATOM	1622	CG	PHE	295	49.439	59.770	24.852	1.00	44.54
	ATOM	1623	CDI	PHE	295	50.325	60.433	25.679	1.00	45.58
	MOTA	1624	CD2	PHE	295	48.853	60.419	23.779	1.00	48.03
35	MOTA	1625	CEI	PHE	295	50.559	61.777	25.480	1.00	49.71
33	ATOM	1626	CEZ	PHE	295	49.086	G1.764	23.583	1.00	47.03
	ATOM	1627	CZ	PHE	295	49.928	62.441	24.444	1.00	52.74
	ATOM ATOM	1628	C	PHE	295	47.765	59.359	27.208	1.00	46.22
	ATOM	1629 1630	O N	PHE	295	47.343	60.503	27.145	1.00	42.87
40	ATOM	1631	Н	LYS	296	48.314	58.905	28.316	1.00	57.30
	ATOM	1632	CA CA	LYS LYS	296	48.708	58.012	28.341	1.00	0.00
		1633	CB	LYS	296 206	48.421	59.737	29.496	1.00	64.18
	ATOM		CG	LYS	296 296	48.497	58.862	30.739	1.00	65.20
	ATOM		CD	LYS	296	48.184	59.668	31.988	1.00	68.41
45		1636	CE	LYS	296	47.966 49.201	58.760	33.189	1.00	72.60
		1637	NZ	LYS	296	50.320	57.947 48.836	33.534	1.00	75.23
	ATOM	1638	HZ	LYS	296	50.088	58.836 59.472	33.780 34.569	1.00	81.49
	ATOM	1639	HZ2	LYS	296	50.511	59.398	32.925	1.00 1.00	0.00
	ATOM	1640	HZ3	LYS	296	51,159	58.270	34.015	1.00	0.00 0.00
50	ATOM	1641	С	LYS	296	49.684	60.547	29.397	1.00	68.26
	ATOM	1542	0	LYS	296	50.729	59.951	29.161	1.00	72.09
	ATOM	1543	N	ASP	297	49.591	61.860	29.569	1.00	72.66
	ATOM		H	ASP	297	49.170	62.489	28.934	1.00	0.00
		1645	CA	ASP	297	50.160	62.460	30.750	1.00	82.26
55	ATOM		CB	ASP	297	51.698	62.430	30.748	1.00	88.06
	MOTA		CG	ASP	297	52.196	61.867	32.082	1.00	92.95
		1648	ODI	ASP	297	51.766	62.345	33.140	1.00	94.73
	ATOM	1649	OD2	ASP	297	53.019	60.945	32.067	1.00	96.27
C C	ATOM		C	ASP	297	49.705	63.893	30.706	1.00	84.91
60		1651	0	ASP	297	49.463	64.419	29.608	1.00	87.39
	MOTA		OT	ASP	297	49.571	64.477	31.777	1.00	84.32
	MOTA		CB	ALA	317	65.517	45.642	-31.211	1.00	60.98
	MOTA		C	ALA	317	63.053	45.718	-30.743	1.00	55.95
	MOTA	1033	0	ALA	317	62.644	46.858	-30.593	1.00	55.33

											0.00
	. 5014	1686	HTI	AL.		317	63.898	47.390	-32.203	1.00	0.00
	MOTA	1656 1657	HT2	AL		317	62.879	46.419	-33.062	1.00 1.00	63.55
	MOTA	1658	N	AL		317	63.865	46.499	-32.753	1.00	0.00
	MOTA	1659	нтз	AL.		317	64.541	46.464	-33.540	1.00	59.35
5	ATOM	1660	CA	AL		317	64.124	45.449	-31,790	1.00	52.67
_	MOTA	1661	N	IL	Ε	318	62.584	44.702	-30.035 -30.235	1.00	0.00
	ATOM	1662	H	11	E	318	62.850	43.777	-28.863	1.00	48.35
	ATOM	1663	CA	11.		318	61.755	44.926 43.749	-28.729	1.00	43.28
	ATOM	1554	CB	п		318	60.789	43.749	-29.796	1.00	39.84
10	ATOM	1665	CG2	. п		318	59.718	42.407	-28.849	1.00	35.21
	ATOM	1666	CG1		.E	318	61.523 60.596	41.194	-28.946	1.00	36.89
	ATOM	1667	CDI		Æ	318	62.702	45.020	-27.656	1.00	48.70
	ATOM		C		Æ	318 318	63.754	44.390	-27.651	1.00	46.83
	ATOM		0		LE	319	62.351	45.808	-26.630	1.00	47.06
15		1670	N		YS YS	319	61.527	46.341	-26.691	1.00	0.00
	ATOM		H		15 YS	319	63.102	45.893	-25.383	1.00	42.30
	ATOM		CA		YS	319	63.641	47,280	-25.184	1.00	46.07
	ATOM		CB CG		YS	319	65.100	47.492	-25.528	1.00	59.38 73.60
	ATOM		CD		YS	319	65.488	48.878	-24.982	1.00 1.00	83.83
2,0	ATON		CE		.YS	319	66.995	49.115	-24,757	1.00	85.82
	MOTA KOTA		NZ		.YS	319	67.263	50.131	-23.739	1.00	0.00
	MOTA		HZ		YS	319	66.878	49.820	-22.825	1.00	0.00
	ATON		HZ	_	.YS	319	66.840	51.037	-24.022	1.00	0.00
25	ATON		HZ		.YS	319	68.294	50.246	-23,654 -24,229	1.00	36.54
23	ATO		С		LYS	319	62.157	45.597	-24.324	1.00	36.02
	ATO		_		LYS	319	60.960	45.806	-23.105	1.00	34.11
	ATO		N		LYS	320	62.635	45.096 44.830	-23.064	1.00	0.00
	ATO		H		LYS	320	63.577	44.993	-21.894	1.00	35.41
30	ATO				LYS	320	61.823 62.389	43.928	-20.978	1.00	36.15
	ATO				LYS	320 320	62.391	42.526	-21.549	1.00	40.56
	ATO				LYS LYS	320 320	63.120	41.594	-20.583	1.00	40.42
	ATO				LYS	320	63.108	40.150	-21.065	1.00	43.28 52.04
	ATO				LYS	320	64.002	39.368	-20.235	1.00	0.00
35					LYS	320	63.689	39.403	-19.245	1.00	0.00
	ATC				LYS	320	64. 96 6	39.751	-20.319	1.00 1.00	0.00
	OTA OTA		-		LYS	320	63.995	38.382	-20.571	1.00	33.67
	ATO				LYS	320	61.701	46.280	-21.053	1.00	31.28
40			_		LYS	320	62.635	47.066		1.00	28.86
-2 (ATO		-		ALA	321	60.521	46.503		1.00	0.00
	AT				ALA.	321	59.777	45.870 47.641		1.00	26.84
	AT		98 C	A	ALA	321	60.307	48.636		1.00	26.05
		OM 16	99 C	В	ALA	321	59.387	47.109		1.00	25.85
4	S AT	OM 17			ALA	321	59.653 58.914			1.00	29.15
		OM 17			ALA	321	59.889			1.00	25.43
	TA	OM 17			HIS	322	60.599				0.00
		OM 17			HIS	322 322	59.185			1.00	20.62
				CA.	HIS HIS	. 322	59.573		6 -14.672		18.17
5				CB	HIS	322	61.001		2 -14.197	1.00	16.95 13.40
	A ⁻	IOM 1		CD2	HIS	322			9 -13.586	1.00	20.74
	N.	TOM 1		ND1	HIS	322		6 48.84	5 -14.409	1.00	0.00
	A.	rom 1	700	HDI	HIS	322	61.98				17.22
_	A	TOM I	709	CEI	HIS	322		2 48.25			19.50
5	55 A	TOM 1	710	NE2	HIS	322					0.00
	^	TOM I	717	HE2	HIS	322	63.43				23.43
	Ą	I MOT		C	HIS	327	2 57.74				
		TOM I		Ö	HIS	32	2 57.44				
			1715	N	ILE	32					
,			1716	H	ILE	32					
			1717	CA	ILE	32			- ·		
			1718	СВ	ILE	32			• • • • • • • • • • • • • • • • • • • •		
		ATOM		CG2	ILE	32	3 54.5	31 43	**************************************		

- 69 -

	17014 17								
	ATOM 17			323	53.411	45.795	-16.461	1.00	17.37
	ATOM 17		ILE ILE	323 323	52.688	44.478	-16.349	1.00	24.35
	ATOM 17		ILE	323	54.891 53.998	48.183 48.930	-15.046	1.00	27.83
5	ATOM 17		GLU	324	55.418	48.357	-15.420 -13.843	1.00 1.00	26.39
	ATOM 17		GLU	324	56.140	47.768	-13.556	1.00	27.46 0.00
	ATOM 17		GLU	324	54.973	49.441	-12.974	1.00	26.27
	ATOM 17:		GLU	324	54.186	48.824	-11.821	1.00	25.82
10	ATOM 17		GLU	324 324	53.692	49.837	-10.816	1.00	23.38
	ATOM 17		GLU	324 324	52.881 52.275	49.122	-9.775	1.00	24.43
	ATOM 17	31 OE2	GLU	324	52.858	48.099 49.586	-10.071 -8.649	1.00	25.59
	ATOM 17		GLU	324	56.191	50.205	-12.459	1.00 1.00	25.24 29.36
15	ATOM 17	-	GLU	324	57.106	49.587	-11.922	1.00	26.17
13	ATOM 173		LYS	325	56.245	51.531	-12.604	1.00	26.30
	ATOM 173		LYS LYS	325	55,526	52.005	-13.083	1.00	0.00
	ATOM 173		LYS	325 325	57.371 58.571	52.335	-12.203	1.00	24.24
	ATOM 173		LYS	325	59.855	52.036 52.715	-13.135 -12.663	1.00	19.60
20	ATOM 173		LYS	325	61.087	52.259	-12.663	1.00 1.00	14.81 11.93
	ATOM 174		LYS	325	61.101	52.703	-14.851	1.00	20,28
	ATOM 174		LYS	325	61.146	54.143		1.00	21.10
	ATOM 174		LYS	723	62.010	54.507	-14.531	1.00	0.00
25	ATOM 174		LYS LYS	325	60.311	54.557	-14.522	1.00	0.00
	ATOM 174		LYS	325 325	61.145	54.397	-15.993	1.00	0.00
	ATOM 174		LYS	325	56.861 55.928	53.774 54.033	-12.340	1.00	26.70
	ATOM 174		ASP	326	57.480	54.700	-13.094 -11.589	1.00 1.00	25.97
2.0	ATOM 174		ASP	326	58.191	54.415	-10.974	1.00	25.25 0.00
30	ATOM 174		ASP	326	57.214	56.136	-11.610	1.00	23.30
	ATOM 175 ATOM 175		ASP	326	57,422	56.737	-13.034	1.00	23.42
	ATOM 175		ASP ASP	326	58.721	56.273	-13.707	1.00	24.74
	ATOM 175		ASP	326 3 26	59.806	56.575	-13.224	1.00	24.42
35	ATOM 175	4 C	ASP	326	58.646 55.835	55.579 56.529	-14.7[]	1.00	21.44
	ATOM 175	5 0	ASP	326	55.284	57.559	-11.122 -11.488	1.00 1.00	20.53 24.45
	ATOM 175		PHE	327	55,222	55.721	-10.264	1.00	19.21
	ATOM 175		PHE	327	55.626	54.862	-10.024	1.00	0.00
40	ATOM 175		PHE	327	53.972	56.107	-9.608	1.00	18.83
	ATOM 175		PHE PHE	327	53.008	54.938	-9.410	1.00	16.54
	ATOM 176		PHE	327 327	52.290 51.070	54.444	-10.658	1.00	20.45
	ATOM 176		PHE	327	52.772	55.004 53.338	-11.016 -11.361	1.00 1.00	17.51
	ATOM 176		PHE	327	50.334	54.424	-12.043	1.00	21.65 20.17
45	ATOM 176		PHE	327	52.022	52.752	-12.380	1.00	18.97
	ATOM 176		PHE	327	50.801	53.298	-12.718	1.00	20.53
	ATOM 176		PHE	327	54.219	56.643	-8.201	1.00	18.93
	ATOM 176		PHE ILE	327 328	55.237	56.323	-7.588	1.00	17.66
50	ATOM 176		ILE	· · 328	53.302 52.602	57.460 57.853	-7.673	1.00	18.15
	ATOM 1770		ILE	328	53.255	57.833 57.771	-8.243 -6.266	1.00	0.00
	ATOM 177		ILE.	328	54.123	59.032	-5.914	1.00	16.96 17.42
	ATOM 177		ILE.	328	53.663	60.311	-6.612	1.00	19.49
55	ATOM 177		ILE	328	54.026	59.244	4.395	1.00	15.51
در	ATOM 1774 ATOM 1775		ILE	328	55.177	60.055	-3.780	1.00	15.10
	ATOM 177		ILE ILE	328	51.802	58.030	-5.976	1.00	18.38
	ATOM 177		ALA	328 329	51.153 51.225	58.692 57.543	-6.769	1.00	18.62
	ATOM 1778		ALA	329	51.223 51.716	57.543 56.924	-4.873 -4.289	1.00	20.05
60	ATOM 1779		ALA	329	49.867	57.908	4.507	1.00 1.00	0.00 20.56
•	ATOM 1780		ALA	329	48.985	56.671	-4.395	1.00	17.55
	ATOM 178		ALA	329	49.854	58.614	-3.168	1.00	24.14
	ATOM 1783 ATOM 1783		YLA	329	50.648	58.289	-2.296	1.00	22.82
	A10M 1/8.	3 N	PHE	330	48.969	59.584	-2.967	1.00	22.84

								1.00	0.00
	ATOM 1784	н	PHE	330	48.352	59.823	-3.6 9 4	1.00	18.35
	ATOM 1785	CA	PHE	330	48.869	60.291	-1.709 -1.937	1.00	18.09
	ATOM 1786	СВ	PHE	330	49.268	61.745	-0.647	1.00	22.46
	ATOM 1787	CG .	PHE	330	49.659	62.459	-0.652	1.00	21.17
5	ATOM 1788	CD1	PHE	330	49.771	63.841 61.760	0.521	1.00	20.68
_	ATOM 1789	CD2	PHE	330	49.931	64.514	0.490	1.00	22.56
	ATOM 1790	CE1	PHE	330	50.147	62.438	1.665	1.00	20.88
	ATOM 1791	CE2	PHE	330	50.309	63.815	1.653	1.00	23.44
	ATOM 1792	CZ	PHE	330	50.409	60.183	-1.260	1.00	20.07
10	ATOM 1793	С	PHE	330	47.425 46.536	60.673	-1.956	1.00	16.75
	ATOM 1794	0	PHE	330	47.169	59.539	-0.117	1.00	20.04
	ATOM 1795	N	CYS	331	47.914	59.170	0.402	1.00	0.00
	ATOM 1796	H	CYS	33 l 33 l	45.831	59.362	0.402	1.00	17.86
	ATOM 1797		CYS	331	45.584	57.950	0.841	1.00	16.56
15	ATOM 1798		CYS	331	45.613	56.747	-0.490	1.00	23.49
	ATOM 1799		CYS CYS	331	45.649	60.235	1.609	1.00	21.19
	ATOM 1800			331	46.586	60.508	2.342	1.00	20.65
	ATOM 1801		CYS SER	332	44.421	60.692	1.836	1.00	22.22
	ATOM 1802		SER	332	43.659	60.414	1.268	1.00	0.00
20	ATOM 1803		SER	332	44.084	61.598	2.928	1.00	21.44
	ATOM 180:		SER	332	42.690	62.162	2.616	1.00	19.68 24.42
			SER	332	41.835	61.127	2.164	1.00	0.00
			SER	332	40.977	61.499	2.030	1.00	21.99
25	ATOM 180 ATOM 180		SER	332	44.120	61.144	4.389	1.00	20.58
25	ATOM 180		SER	332	44.290	61.947	5.302	1.00 1.00	23.27
	ATOM 181		SER	333	43.945	59.844	4.657	1.00	0.00
	ATOM 181		SER	333	43.779	59.215	3.918	1.00	20.50
	ATOM 181		SER	333	43.971	59.274	5.984 6.523	1.00	21.15
30	ATOM 181		SER	333	42.580	58.895		1.00	23.77
50	ATOM 18		SER	333	41.566	58.641	5.556 6.021	1.00	0.00
	ATOM 18		SER	333	40.748	58.468	5.819	1.00	23.49
	81 MOTA		SER	333	44.745	57.979	4.736	1.00	19.23
	ATOM 18	17 0	SER	333	45.104	57.510	6.951	1.00	23.40
35	ATOM 18	18 N	THR	334	45.003	57.359 57.711	7.788	1.00	0.00
	ATOM 18		THR	334	44.646	56.099	6.997	1.00	24.09
	ATOM 18		THR	334	45.720	56.407	8.296	1.00	28.30
	ATOM 18		THR	334	46.501 47.797	55.855	8.169	1.00	34.11
		22 OG1	THR	334	48.191	56.183	7.351	1.00	0.00
40		323 HG1	THR	334 334	45.753	55.956	9.519	1.00	14.09
		324 CG2	THR	334	44.639	54.990	6.904	1.00	22.20
		325 C	THR	334	43.459	55.273	7,098	1.00	23.69
		826 O	THR PRO	335	44.853	53.756	6.619	1.00	20.97
		827 N	PRO	335	46.175	53.207	6.414	1.00	15.82
4			PRO	335	43.804	52.736	6.469	1.00	19.65
	ATOM 1		PRO	335	44.565	51.473	6.155	1.00	20.20 20.66
			PRO	335	45.823	52.030	5.529	1.00	25.53
	• • • • •		PRO	335	42.850	52.545	7.636	1.00	29.15
_	• • • • •	.832 C 1833 O	PRO	335	43.309	52.608	8.769	1.00	24.20
5	MOTA 0		ASP	336	41.545	52.314	7.406	1.00 1.00	0.00
	ATOM		ASP	336	41.220	52.278	6.479	1.00	28.39
	ATOM	1836 CA	ASP	336	40.529	52.137	8.449 9,710	1.00	32.12
	MOTA		ASP	336	40.940		9.710	1.00	43.11
5	5 ATOM		ASP	336	41.344		10.040	1.00	52.99
ر	MOTA	1839 OD		336	42.285		8.531	1.00	51.49
		1840 OD		336	40.725		9.046	1.00	29.38
	MOTA	1841 C	ASP	336				1.00	34.19
	MOTA		ASP	336				1.00	25.69
-	MOTA 0	1843 N	ASN	337				1.00	0.00
		1844 H	ASN	337				1.00	23.68
	MOTA			337					22.73
	ATOM	1846 CE	asn.	337					22.82
		1847 CC	ASN .	337	42.06	5 55.421	22.000	2.24	

- 71 -

	ATOM 18	48 OD1	ASN	337	41.379	£4.601 ·			
	81 MOTA		ASN	337	43.293	54.991 54.940	11.937	1.00	25.91
	ATOM 18:		ASN	337	43.822	55.204	10.989 10.218	1.00	18.06
5	ATOM 185		ASN	337	43.604	54.320	11.691	1.00	0.00
5		-	ASN	337	39.600	56.812	8.496	1.00 1.00	0.00
	ATOM 185 ATOM 185		ASN	337	39,737	56.860	7.283	1.00	25.85 29.96
			VAL	338	38.815	57.691	9.095	1.00	27,31
	ATOM 185	•	VAL	338	38.760	57.662	10.072	1.00	0.00
10	ATOM 185		VAL	338	38.117	58.722	8.361	1.00	24.45
	ATOM 185		VAL	338	36.902	59.297	9.168	1.00	24.69
	ATOM 185		VAL	338	35.901	58.187	9.486	1.00	25.15
	ATOM 186		VAL VAL	338	37.369	59.946	10.467	1.00	25.43
	ATOM 186	-	VAL	338	39.045	59.870	8.024	1.00	25.97
15	ATOM 186	_	SER	338 339	40.124	60.040	8.588	1.00	26.18
	ATOM 186		SER	339	38.588	60.680	7.067	1.00	27.49
	ATOM 186	4 CA	SER	339	37.756 39.231	60.450	6.596	1.00	0.00
	ATOM 186	5 CB	SER	339	39.666	61.948	6.743	1.00	32.77
	ATOM 186	6 OG	SER	339	39.874	62.030 60.771	5.285	1.00	30.77
20	ATOM 186	7 HG	SER	339	39.097	60.209	4.672	1.00	38.35
	ATOM 186	-	SER	339	38.181	63.015	4.677	1.00	0.00
	ATOM 1869	_	SER	339	37.008	62.669	6.948 6.924	1.00	32.35
	ATOM 1870		TRP	340	38.549	64.280	7.130	1.00 1.00	29.99
25	ATOM 187		TRP	340	39,494	64_546	7.074	1.00	31.11 0.00
23	ATOM 1872 ATOM 1872		TRP	340	37.580	65.306	7,447	1.00	32.42
	ATOM 1873 ATOM 1874		TRP	340	38.110	66.071	8.665	1.00	28.07
	ATOM 187		TRP	340	38.069	65.153	9.851	1.00	31.22
	ATOM 1876		TRP TRP	340	36.790	64.901	10.522	1.00	33.69
30	ATOM 1877	CE3	TRP	340	37.281	63.820	11.450	1.00	38.45
	ATOM 1878	CDI	TRP	340 340	35.473 39.112	65.302	10.526	1.00	37.14
	ATOM 1879		TRP	340	38.611	64.394	10.320	1.00	32.03
	ATOM 1880	HEI	TRP	340	39.143	63.616	11.256	1.00	35.26
2-	ATOM 1881	CZ2	TRP	340	36.364	62.912 63.217	11.687	1.00	0.00
35	ATOM 1882		TRP	340	34.612	64.660	12.277 11.389	1.00	43.31
	ATOM 1883		TRP	340	35.043	63.639	12.227	1.00 1.00	42.02 45.40
	ATOM 1884	•	TRP	340	37.260	66.265	6.311	1.00	35.22
	ATOM 1885 ATOM 1886		TRP	340	38.137	66.652	5.544	1.00	36.37
40	ATOM 1887	•	ARG	341	35.989	66.670	6.210	1.00	37.03
	ATOM 1888		ARG	341	35.321	66.289	6.821	1.00	0.00
	ATOM 1889		ARG ARG	341	35.487	67.607	5.211	1.00	36.96
	ATOM 1890	CG	ARG	341 341	34.687	66.815	4.170	1.00	34.84
	ATOM 1891		ARG	341	34.391	67.533	2.861	1.00	37.33
45	ATOM 1892	NE	ARG	341	33.517 32.151	66.655	1.972	1.00	39.02
	ATOM 1893		ARG	341	31.684	66.768 67.621	2.441 2.324	1.00	48.98
	ATOM 1894		ARG	341	31.501	65.767	3.031	1.00 1.00	0.00
	ATOM 1895		ARG	341	32.089	64.556	3.232	1.00	51.01 56.13
50	ATOM 1896		ARG	341	33.030	64.400	2.934	1.00	0.00
J 0	ATOM 1897 ATOM 1898		ARG	** 341	31.577	63.821	3.676	1.00	0.00
		NH2	ARG	341	30.219	65.962	3.440	1.00	58.28
	ATOM 1899 ATOM 1900		ARG	341	29.777	66.848	3.294	1.00	0.00
	ATOM 1901	HH22 C	ARG	341	29.720	65.218	3.884	1.00	0.00
55	ATOM 1902	Ö	ARG ARG	341	34.616	68.684	5.864	1.00	37.23
	ATOM 1903	Й	HIS	34l	33.969	68.486	6.883	1.00	38.61
	ATOM 1904	H	HIS	342 342	34.550 35.002	69.880	5.324	1.00	39.36
	ATOM 1905	CA	HIS	342	33.820	70.055	4,471	1.00	0.00
	ATOM 1906	CB	HIS	342	34.885	70.966 71.921	5.934	1.00	43.08
60	ATOM 1907	CG	HIS	342	34.284	73.095	6.463 7.199	1.00 1.00	41.90 45.95
	ATOM 1908	CD2	HIS	342	33.508	74.035	6.600	1.00	45.95 46.70
	ATOM 1909	NDI	HLS	342	34.247	73.368	8.511	1.00	48.38
	ATOM 1910	HDI	HIS	342	34.661	72.884	9.248	1.00	0.00
	ATOM 1911	CEI	HIS	342	33.458	74.387	8.703	1.00	48.44
						-			70,77

								1.00	49.68
	ATOM 1912	NE2	HIS	342	33.020	74.776	7.542 7.407	1.00	0.00
		HE2	HIS	342	32.497	75.591	4.899		46.18
	ATOM 1913 ATOM 1914	C	HIS	342	32.894	71.612	3.826		43.57
	ATOM 1915	ō	HIS	342	33.367	71.960 71.844	5.101	1.00	48.28
5	ATOM 1916		PRO	343	31.637	71.752	6.398	1.00	46.23
ر	ATOM 1917	CD	PRO	343	30.984	72.297	4.073	1.00	49.91
	ATOM 1918	CA	PRO	343	30.678	72.590	4.824	1.00	44.40
	ATOM 1919	CB	PRO	343	29.387 29.512	71.689	6.015	1.00	41.79
	ATOM 1920	CG	PRO	343	31.068	73.489	3.199	1.00	53.27
10	ATOM 1921	С	PRO	343	30.905	73.585	1.988	1.00	55.64
	ATOM 1922	Ο.	PRO	343 344	31.615	74.459	3.909	1.00	54.13
	ATOM 1923	N	THR	344 344	31.830	74.264	4.837	1.00	0.00
	ATOM 1924	H	THR	344	31.911	75.771	3.354	1.00	53.54
	ATOM 1925	CA	THR	344	31.577	76.802	4.429	1.00	56.09 63.15
15	ATOM 1926	CB	THR THR	344	31,901	76.175	5.672	1.00	0.00
	ATOM 1927	OG!	THR	344	31.164	75.566	5.752	1.00	57.44
	ATOM 1928	HG1	THR	344	30.088	77.164	4.490	1.00 1.00	53.06
	ATOM 1929	CG2 C	THR	344	33.367	75.826	2.944	1.00	53.90
	ATOM 1930	0	THR	344	33.802	76.516	2.046	1.00	51.25
20	ATOM 1931	Ŋ	MET	345	34.192	75.055	3.632	1.00	0.00
	ATOM 1932	H	MET	345	33.844	74.348	4.202 3.363	1.00	52.69
	ATOM 1933 ATOM 1934	Ċλ	MET	345	35.604	75.172	4.662	1.00	60.31
	ATOM 1934 ATOM 1935	CB	MET	345	36.357	75.229	5,498	1.00	71.30
25	ATOM 1936	ČĞ	MET	345	36.128	76.474	7.176	1,00	86.57
25	ATOM 1937	SD	MET	345	36.752	76.178	7.073	1.00	81.62
	ATOM 1938	CE	MET	345	38.405	76.807 74.006	2.553	1.00	47.27
	ATOM 1939	С	MET	345	36.133	74.042	2.179	1.00	48.63
	ATOM 1940	0	MET	345	37.288	72.947	2.236	1.00	40.77
30		N	GLY	346.	35.383	72.892	2.503	1.00	0.00
.50	ATOM 1942	H	GLY	346	34.439	71.822	1.544	1.00	36.57
	ATOM 1943	CA	GLY	346	35.981 36.764	70.915	2.495	1.00	35.51
	ATOM 1944	C	GLY	346	36.680	70.988	3.718	1.00	40.26
	ATOM 1945		GLY	346	37.567	70.012	1.949	1.00	32.41
35	ATOM 1946		SER	347 347	37.693	70.023	0.980	1.00	0.00
	ATOM 1947	-	SER	347	38.274	68.999	2.698	1.00	28.45 24.81
	ATOM 194		SER SER	347	38.454	67.768	1.820	1.00	30.64
	ATOM 194		SER	347	37.262	67.491	1.098	1.00 1.00	0.00
	ATOM 195		SER	347	36.547	67.238	1.697	1.00	31.86
4		•	SER	347	39.631	69.491	3.157	1.00	33.18
	ATOM 195		SER	347	40.397	70.136	2.438	1.00	29.14
	ATOM 195	-	VAL	348	39.920	69.144	4,411 4,887	1.00	0.00
	ATOM 195 ATOM 195		VAL	348	39.306	68.543	5.086	1.00	25.67
4	5 ATOM 19	· ·	VAL	348	41.127	69.588	6.494	1.00	27.65
4	ATOM 19		VAL	348	41.215	68.976	7.309	1.00	29.29
	ATOM 19		VAL	348		69.746 69.060	7.219	1.00	29.60
	ATOM 19		VAL	348	44		4.305	1.00	25.79
	ATOM 19		VAL	348			4.071	1.00	31.26
	0 ATOM 19		VAL	. 348			3.862	1.00	23.67
-	ATOM 19		PHE	349			4.013	1.00	0.00
	ATOM 19	63 H	PHE	349			3.181	1.00	22.69
	ATOM 19	64 CA	PHE	349			2,908	1.00	18.76
		965 CB	PHE	349			1.941		20.55
	55 ATOM 1	966 CG	PHE	349			0.702	1.00	14.77
•		967 CD		34					15.53
		968 CD		34		*			16.44
	ATOM 1	969 CE		34			- 440		13.87
	ATOM 1	970 CE		34	·				
		1971 CZ		34					
		1972 C	PHE	34	19 44.96	484	1.609		
	MOTA	1973 0	PHE			40.40			
		1974 N	ILE		50 42.79 50 41.9			6 1.00	0.00
	MOTA	1975 H	ILE	. 3	30 71.3				

- 73 -

	MOTA	1976	~.							
		1977	CA CB	ILE ILE	350	42.969	69.170	-0.185	1.00	23.11
		1978	CG2	ILE	350 350	41.617 41.741	69.131	-0.939	1.00	23.98
		1979	CGI	ILE	350		59.922 67.689	-2.229	1.00	24.02
5	ATOM		CDI	ILE	350	42.089	66.972	-1.267 -2.272	00.1 00.1	20.37
	ATOM		С	IL.E	350	43.451	70.600	0.073	1.00	14.33 23.87
	ATOM		0	ILE	350	44.428	71.058	-0.510	1.00	25.61
	ATOM		. N	GLY	351	42.813	71.359	0.962	1.00	21.19
10	ATOM		H	GLY	351	42.057	70.988	1.467	1.00	0.00
10		1985	CA	GLY	351	43.248	72.720	1.212	1.00	19.54
		1986 1987	C 0	GLY	351	44.702	72.800	1.640	1.00	26.25
	ATOM		N	GLY ARG	351	45.524	73.568	1.138	1.00	28.41
	ATOM		H	ARG	352 352	45.064	71.963	2.614	1.00	28.53
15	ATOM		CA	ARG	352 352	44.388 46.449	71.391	3.045	1.00	0.00
	ATOM		CB	ARG	352	46.589	71.872 70.842	3.042	1.00	26.92
	ATOM :	1992	CG	ARG	352	47.697	71.286	4.167 5.104	1.00 1.00	31.79
	ATOM		CD	ARG	352	47.091	71.997	6.310	1.00	35.13 38.95
•	MOTA		NE	ARG	352	47.179	71.107	7.452	1.00	44.61
2,0	ATOM		HE	ARG	352	47.920	70.468	7.491	1.00	0.00
		1996	CZ	ARG	352	46.307	71.116	8.449	1.00	41.51
	ATOM 1		NH1	ARG	352	46.502	70.272	9.494	1.00	48.56
		1999	HH11 HH12	ARG	352	47.294	69.660	9.507	1.00	0.00
25		2000	NH2	ARG ARG	352 352	45.849	70.267	10.251	1.00	0.00
		2001	HH21	ARG	352	45.227 45.080	71.943	8.435	1.00	44.65
		2002	HH22	ARG	352	44.570	72.554 71.936	7.657	1.00	0.00
	ATOM 2	2003	С	ARG	352	47.419	71.483	9.187 1.931	1.00	0.00
	ATOM 2	2004	0	ARG	352	48.535	71.995	1.866	1.00	26.87 26.27
30		2005	N	LEU	353	47.033	70.567	1.037	1.00	25.01
	ATOM 2		Н	LEU	353	46.156	70.135	1.143	1.00	0.00
	ATOM 2		CA	LEU	353	47.866	70.154	-0.070	1.00	23.26
	ATOM 2		CB	LEU	353	47.222	68. 94 7	-0.778	1.00	25.55
35	ATOM 2		CG CD1	LEU	353	47.975	68.452	-2.020	1.00	23.59
• •	ATOM 2		CD2	LEU LEU	353 353	49.420	68.121	-1.696	1.00	25.86
	ATOM 2		C	LEU	353	47.258 48.034	67.230 71.303	-2.552	1.00	29.54
	ATOM 2		ō	LEU	353	49.142	71.503	-1.031 -1.485	1.00	27.31
	ATOM 2	014	N	ILE	354	46.923	71.977	-1.463	1.00 1.00	24.83 22.68
40	ATOM 2		H	ILE	354	46.049	71.660	-1.035	1.00	0.00
		016	CA	ILE	354	47.019	73.174	-2.168	1.00	24.25
		017	CB	ILE	354	45.595	73.782	-2.327	1.00	21.11
	ATOM 2		CG2	ILE	354	45.652	75.176	-2 .940	1.00	21.96
45	ATOM 2		CG1	ILE	354	44.769	72.893	-3.248	1.00	17.07
13	ATOM 2		CD1	ILE	354	43.279	73.267	-3.187	1.00	18.04
	ATOM 2		0	ile Ile	354 354	48.000	74.172	-1.540	1.00	25.12
	ATOM 2		N	GLU	355	48. 96 7 47.806	74.616 74.559	-2.165	1.00	27.26
	ATOM 2		H	GLU	355	47,040	74.212	-0.276 0.235	1.00 1.00	27.20
50	ATOM 2	025	CA	GLU	355	48.719	75.507	0.233	1.00	0.00 25.06
	ATOM 2		CB	GLU	355	48.501	75.685	1.803	1.00	32.73
	ATOM 2		CG	GLU	355	47.165	76.212	2.282	1.00	45.70
	ATOM 2		CD	GLU	355	47.053	75.777	3.746	1.00	58.26
55	ATOM 2		OEI	GLU	355	45.921	75.683	4.245	1.00	61.90
33	ATOM 2		OE2	GLU	355	48.092	75.513	4.385	1.00	61.89
	ATOM 2		C	GLU	355	50.150	75.114	0.242	1.00	23.42
	ATOM 2		0 N	GLU	355	51.029	75.893	-0.054	1.00	25.26
	ATOM 2		H	HIS HIS	356 356	50.461	73.862	0.489	1.00	26.55
60	ATOM 2		CA	HIS	356 356	49.739	73.225	0.663	1.00	0.00
	ATOM 2		CB	HIS	356	51.855 51.997	73.438	0.432	1.00	26.58
	ATOM 2		CG	HIS	356	51.846	72.019 72.065	1.041	1.00	27.11
	ATOM 2		CD2	HIS	356	52.890	72.063 72.264	2.551 3.421	1.00 1.00	23.74
	ATOM 2		NDI	HIS	356	50.728	72.044	3.421	1.00	22.80 25.98
				_			78.074	J.200	1.00	43.70

			****	766	49.819	71.907	2.953	1.00	0.00
	ATOM 2040	HDI	HIS	356	51.036	72.231	4.550	1.00	19.00
	ATOM 2041	CEI	HIS	356	52.333	72.360	4.608	1.00	22,63
,	ATOM 2042	NEI .	HIS	356	-	72.450	5.460	1.00	0.00
	ATOM 2043	HE2	HIS	356	52.814	73.447	-0.983	1.00	26.45
5	ATOM 2044	C	HIS	356	52.411	73.809	-1.239	1.00	20.78
_	ATOM 2045	0	HIS	356	53.560	73.039	-1.948	1.00	17.38
	ATOM 2046	N	MET	357	51.586	72.732	-1.747	1.00	0.00
	ATOM 2047	. H	MET	357	50.672		-3.316	1.00	28.28
	ATOM 2048	CA	MET	357	52.037	73.060	4,239	1.00	30.75
10	ATOM 2049	CB	MET	357	50.995	72.413	4.254	1.00	34.53
	ATOM 2050	CG	MET	357	50.860	70.870	-5,054	1,00	43.28
	ATOM 2051	SD	MET	357	52.182	69.919	-6.584	1.00	37.66
	ATOM 2052	CE	MET	357	51.449	69.433	-3.704	1.00	29.20
	ATOM 2053	С	MET	357	52.249	74.503		1.00	31.64
15	ATOM 2054	Ō	MET	357	53.237	74.846	-4.338	1.00	25.95
13	ATOM 2055	N	GLN	358	51.365	75.428	-3.360	1.00	0.00
	ATOM 2056	H	GLN	358	50.539	75.168	-2.897	1.00	27.58
	ATOM 2057	CA	GLN	358	51.650	76.808	-3.709	1.00	24.97
	ATOM 2058	CB	GLN	358	50.505	77.693	-3.214	1.00	26.18
20	ATOM 2059	CG	GLN	358	49.262	77.438	-4.067	1.00	28.54
20	• • • • • • • • • • • • • • • • • • • •	CD	GLN	358	48.063	78.234	-3.596		30.13
		OEI	GLN	358	47.170	78.570	-4.362	1.00	31.54
	ATOM 2061	NE2	GLN	358	47.919	78.558	-2.329	1.00	0.00
	ATOM 2062		GLN	358	48.611	78.285	-1.693	1.00	
	ATOM 2063	HE21	GLN	358	47.124	79.077	-2.095	1.00	0.00
25	ATOM 2064	HE22	GLN	358	52.982	77.344	-3.182	1.00	30.98
	ATOM 2065	C	GLN	358	53.783	77.971	-3.869	1.00	31.45
	ATOM 2066	0		359	53.234	77,083	-1.906	1.00	32.65
	ATOM 2067	N	GLU GLU	359	52.581	76.563	-1.386	1.00	0.00
	ATOM 2068	H		359	54.425	77.558	-1.243	1.00	32.96
30	ATOM 2069	CA	GLU	359	54.178	77.382	0.251	1.00	37.33
	ATOM 2070	CB	GLU	359	55.215	77.874	1.245	1.00	51.69
	ATOM 2071	CG	GLU		55.241	79.379	1.308	1.00	63.61
	ATOM 2072		GLU	359	54.197	79.992	1.550	1.00	70.65
	ATOM 2073		GLU	359		79.940	1.132	1.00	69.55
35	ATOM 2074	OE2	GLU	359	56.320	76.809	-1.721	1.00	33.1 9
	ATOM 2075	С	GLU	359	55.638	77.422	-1.974	1.00	36.20
	ATOM 2076	, 0	GLU .	359	56.664	75.475	-1.867	1.00	33.15
	ATOM 2077	N	TYR	360	55.584	75.002	-1.843	1.00	0.00
	ATOM 2078	H	TYR	360	54.723	74.710	-2.053	1.00	28.52
40		CA CA	TYR	360	55.814	73.528	-1.079	1.00	30.16
	ATOM 2080	CB	TYR	360	56.902		0.333	, 1.00	28.83
	ATOM 208	1 CG	TYR	360	56.910	74.022	1.087	1.00	30.92
	ATOM 208	2 CD1	TYR	360	55.765	73.873	2.342	1.00	33.76
	ATOM 208		TYR	360	55.711	74.433	0.824	1.00	30.38
4 !			TYR	360	58.022	74.677	2.080	1.00	32.36
3.	ATOM 208		TYR	360	57.974	75.239	2.823	1.00	35.09
	ATOM 208		TYR	360	56.813	75.114	4.073	1.00	41.75
	ATOM 208		TYR	360	56.737	75.692	4.430	1.00	0.00
	ATOM 201		TYR	360	55.852	75.603		1.00	27.85
			TYR	360	57.148	74.120	-3.389	1.00	27.93
5	ATOM 20		TYR	360	58.243	73.583	-3.525		27.66
	ATUM 20	90 O	ALA	361	56.288	74.169	-4.411	1.00	0.00
	ATOM 20	92 H	ALA	361	55.459	74.682	-4.320	1.00	28.82
	ATOM 20		ALA	361	56.544	73.494	-5.678	1.00	_
	ATOM 20	93 CA	ALA	361			-6,600	1.00	26.34
5	5 ATOM 20	194 CB	ALA	361			-6.355	1.00	32.26
	ATOM 20	95 C		361			-7.116	1.00	35.01
	ATOM 20	96 O	ALA	362			-6.133	1.00	38.50
	ATOM 20	797 N	CYS	362			-5.620	1.00	0.00
	ATOM 20	14 8ec	CYS				-6.659	1.00	42.41
•	60 ATOM 2	099 CA		362		~		4	41.51
	ATOM 2	100 C	CYS	363		_, _,			41.46
	ATOM 2	101 0	CYS	36					
	ATOM 2	102 CB		36					
	ATOM 2	103 SG	CYS	36	2 61.40	U 11.033	-,.050		

- 75 -

	ATOM		N	SER	363	60.741	74.757	-4.681	1.00	40.95
	ATOM		H	SER	363	59.909	74.803	-4.165	1.00	0.00
	ATOM		CA	SER	363	61.955	74.338	-4.026	1.00	43.35
5	ATOM		CB	SER	363	62.122	75.236	-2.800	1.00	44.65
5	MOTA		OG	SER	363	60.899	75.837	-2.362	1.00	53.93
	ATOM		HG	SER	363	60,553	76.414	-3.046	1.00	0.00
	ATOM		C	SER	363	61.985	72.866	-3.661	1.00	40.77
	ATOM ATOM		0	SER	363	63.036	72. 2 55	-3.593	1.00	43,37
10	MOTA		N H	CYS	364	60.857	72.220	-3.399	1.00	38.31
	MOTA		CA	CYS CYS	364	59.976	72.633	-3.542	1.00	0.00
	ATOM		CB	CYS	364	60.937	70.839	-2.979	1.00	33.39
	ATOM		SG	CYS	364 364	60.056	70.623	-1.755	1.00	32.55
	ATOM		C	CYS	364	60.508 60.478	71.767 69.967	-0.422	1.00	40.02
15	ATOM		ō	CYS	364	59.610	70.346	-4.121 -4.887	1.00	29.84
	ATOM		N	ASP	365	61.025	68.770	-4.300	1.00	30.07 28.01
	ATOM	2120	H	ASP	365	61.859	68.579	-3.828	1.00	0.00
	ATOM	2121	CA	ASP	365	60.406	67.784	-5.164	1.00	27.07
	ATOM		CB	ASP	365	61.379	66.619	-5.419	1.00	27.73
20	ATOM		CG	ASP	365	61.757	65.825	-4.171	1.00	31.83
	ATOM		ODI	ASP	365	61.026	64.918	-3.783	1.00	34.71
	ATOM		OD2	ASP	365	62.805	66.094	-3.599	1.00	42.02
	ATOM		С	ASP	365	59.111	67.254	-4.538	1.00	28.95
25	MOTA		0	ASP	365	58.838	67.420	-3.343	1.00	27.98
25	ATOM ATOM		N	VAL	366	58.277	66.590	-5.336	1.00	27,55
	ATOM		H	VAL	366	58.544	66.434	-6.268	1.00	0.00
	ATOM		CA CB	VAL	366	56.968	66.183	-4.857	1.00	27.84
	MOTA		CGI	VAL VAL	366	56.251	65.513	-6.051	1.00	24.13
30	ATOM		CG2	VAL	366	55.102	64.601	-5.672	1.00	23.78
	ATOM		C	VAL	366 366	55.629 57.007	66.654	-6.848	1.00	23.21
	ATOM		ŏ	VAL	366	56.256	65.309 65.336	-3.611	1.00	28.22
	ATOM		N	GLU	367	57.856	64.298	-2.674 -3.489	1.00 1.00	30.81 28.49
	ATOM	2137	н	GLU	367	58.450	64.111	-4.237	1.00	0.00
35	ATOM	2138	CA	GLU	367	57.867	63.516	-2.259	1.00	27.85
	ATOM		CB	GLU	367	58.939	62.428	-2.289	1.00	26.64
	MOTA		CG	GLU	367	58.553	61.219	-3.144	1.00	28.69
	ATOM		CD	GLU	367	59.442	60.019	-2.820	1.00	37.41
4.0	ATOM		OEI	GLU	367	59.643	59.736	-1.632	1.00	42.87
40	MOTA		OE2	GLU	367	59.937	59.388	-3.767	1.00	37.88
	MOTA MOTA		C ,		367	58.106	64.352	-1.012	1.00	28.61
	MOTA		O N	GLU GLU	367	57.604	64.052	0.059	1.00	31.13
	ATOM		H	GLU	368 368	58.872	65.437	-1.099	1.00	28.39
45	ATOM		CA	GLU	368	59.242 59.094	65.725 66.231	-1.959 0.076	1.00	0.00
	ATOM		CB	GLU	368	60.278	67.145	0.076 -0.069	1.00 1.00	26.65
	ATOM		CG	GLU	368	61.017	66.888	1.244	1.00	34.26 53.59
	ATOM		CD	GLU	368	61.471	68.195	1.841	1.00	63.52
	ATOM	2152	OEI	GLU	368	61.848	69.099	1.078	1.00	71.58
50	MOTA	2153	OE2	GLU	368	61.444	68,298	3.071	1.00	64.40
	MOTA	2154	C	GLU	368	57.923	67.082	0.396	1.00	23.42
	ATOM		0	GLU	368	57.561	67.224	1.555	1.00	27.18
	ATOM		N	ILE	369	57.268	67.677	-0.591	1.00	22.22
	ATOM		Н	ILE	369	57.595	67.596	-1.513	1.00	0.00
55	ATOM		CA	ILE.	369	56.027	68.395	-0.295	1.00	19.89
	ATOM		CB	ILE	369	55.403	68.931	-1.624	1.00	22.54
	ATOM		CG2	ILE	369	54.089	69.671	-1.355	1.00	20.28
	MOTA		CGI	ILE	369	56.360	69.917	-2.281	1.00	17.77
60	MOTA MOTA		CDI	ILE	369	55.911	70.256	-3.701	1.00	15.38
	ATOM		0	ILE ILE	369 360	55.024	67.497	0.435	1.00	20.18
	MOTA		N	PHE	369 370	54.331	67.885	1.371	1.00	20.19
	ATOM		H	PHE	370 370	54.923 55.470	66.241	0.020	1.00	21.39
	ATOM		CA	PHE	370	55.470 53.962	65.932 65.359	-0.733 0.654	1.00	0.00 23.30
					210	33.704	27.773	0.034	1.00	23.30

							44 MD.	-0.121	1.00	23.30
	ATOM	2168	СВ	PHE	370	53.842	64.029 64.222	-1.451	1.00	23.92
	ATOM	2159	CG	PHE	370	53.140	63.153	-2.304	1.00	27.01
	ATOM		CDI	PHE	370	53.037	65,444	-1.841	1.00	29.88
	ATOM		CD2	PHE	370	52.624	63.311	-3.533	1:.00	26.34
5	ATOM	2172	CEI	PHE	370	52.427 52.013	65.595	-3.069	1.00	27.67
	ATOM	2173	CEI	PHE	370	51.906	64.521	-3,919	1.00	27.89
	ATOM	2174	ÇΖ	PHE	370	54.360	65.076	2.069	1.00	23.38
	ATOM		. C	PHE	370 370	53.543	65.031	2.978	1.00	24.64
	ATOM		0	PHE ARG	371	55.654	64.883	2.314	1.00	24.62 0.00
10	ATOM		N U	ARG	371	56.291	64.804	1.573	1.00	26.21
	ATOM		H. CA	ARG	371	56.103	64.783	3.683	1.00	25.42
	ATOM	1 2179 1 2180	CB	ARG	371	57.611	64.627	3.722	1.00 1.00	26.24
	ATON	2181	CG	ARG	371	5 7.954	64.338	5.180	1.00	27.46
15	ATOL	1 2182	CD	ARG	371	59.453	64.358	5.411	1.00	34.21
7.3	ATON	1 2183	NE	ARG	371	60.022	65.650	5.122 4,286	1.00	0.00
	ATO!	1 2184	HE	ARG	371	60.521	65.766	5.969	1.00	40.22
	ATON	4 2185	CZ	ARG	371	59.899	66.694	7.152	1.00	39.64
	ATO	1 2186	NHI	ARG	371	59.207	66.648 65.805	7.435	1.00	0.00
20	ATO	4 2187	HHII	ARG	371	58.750	67.463	7.729	1.00	0.00
	ATO!	M 2188	HH12	ARG	371	59.163 60.525	67.849	5.632	1.00	39.43
	ATO	M 2189	NH2	ARG	371	61.067	67.884	4.795	1.00	0.00
	ATO	M 2190	HH21	ARG	371 371	60.471	68.649	6.231	1.00	0.00
	OTA	M 2191	HH22		371	55.704	65.999	4.505	1.00	29.69
25	ATO	M 2192	c	ARG ARG	371	55.105	65.907	5.576	1.00	32.42 26.46
	OTA	M 2193	0	LYS	372	56.020	67.200	4.020	1.00	Q.00
		M 2194		LYS	372	56.525	67.281	3.182	1.00	23.67
	ATC	M 2195	H CA	LYS	372	55.576	68.377	4.733	1.00 1.00	25.18
20	ATC	M 2196 M 2197	CB	LYS	372	56.008	69.609	3.976	1.00	21.18
30	AIC	M 2198	CG	LYS	372	57.506	69.718	4.053	1.00	33.26
	ATO	M 2199		LYS	372	5 7.933	70.852	3.136 3.789	1.00	42.84
	ATO	OM 2200	CE	LYS	372	59.042	71.666	4.969	1.00	50.90
	ATO	OM 220	NZ.	LYS	372	58.499	72.323 71.612	5,636	1.00	0.00
35		OM 220	HZ1		372	58.138	72.944	4.674	1.00	0.00
	AT	OM 220	3 HZ2		372	57.717 59.239	72.893	5.426	1.00	0.00
	AT	OM 220	4 HZ3	LYS	317		68.434	4.960	1.00	22.44
	ΑT	OM 220	5 C	LYS	372 372		68.867	6.008	1.00	25.03
		OM 220		LYS	373		68.015	4.026	1.00	24.85
4(OM 220		VAL VAL	373		67.723	3.154	1.00	0.00 26.14
		OM 220		VAL	373		67.977	4.333	1.00	29.33
		OM 220		VAL	. 373			3.121	1.00 1.00	23.06
	AT.	TOM 22 TOM 22						3.541	1.00	20.17
	E A	TOM 22	12 CG	·		3 51.108		1.933 5.471	1.00	26.25
4	⊃ <u>^</u> .	TOM 22	13 C	VAL		3 51.588		6.336	1.00	25.40
	<u> </u>	TOM 22	14 O	VAL				5.540	1.00	22.91
			15 N	ARG				4.838	1.00	0.00
		TOM 2		ARC				6,660	1.00	24.92
5	0 A	TOM 2	217 CA					6.511	1.00	24.92
_	A	TOM 2	218 CE					5.307	1.00	26.50
		TOM 2	219 CC			74 52.82		5.253	1.00	21.69
		TOM 2	220 CI			74 54.26		5.278	1.00	21.59
	- 1	TOM 2	221 N			74 54.72			1.00	0.00
5	55 /	TOM 2	222 H		_	74 54.97		4,161		19.13 22.36
	4	ATOM 2	223 C		-	74 54.33				0.00
		ATOM 2			_	74 53.3	40 61.217			0.00
		ATOM 2	225 R			74 54.8				
	- ^	MOTA	7440 N	IH12 AR IH2 AR		374 56.3	44 61.04		·	
	60	ATOM :	1111 N	TH21 AR		374 56.8				
		ATOM :	1110 E	H22 AF		374 56.8				
		MOTA MOTA		AF		374 52.4	44 44			
		ATOM		•		374 51.7	124 65.53	9 6.93		
		YIOM	، دیشم	-						

	ATOM 2232	N	PHE	375	53.581	66.204	7.996	1:00	24.0
	ATOM 2233	H	PHE	375	5 4.147	66.181	7.193	1:00 1:00	24.10
	ATOM 2234	CA	PHE	375	54.025	66.896	9.187	1.00	0.00 23,34
_	ATOM 2235	CB	PHE	375	55.388	67.523	8.828	1.00	29.16
5	ATOM 2236	CG	PHE	375	56.010	68.272	9.989	1.00	35.30
	ATOM 2237	CDI	PHE	375	56.926	67.655	10.803	1.00	36.99
	ATOM 2238	CD2	PHE	375	35.684	69.589	10.221	1.00	33.55
	ATOM 2239	.CEI	PHE	375	57.518	68.348	11.838	1.00	36.47
10	ATOM 2240	CEZ	PHE	375	56.281	70.277	11.255	1.00	38.44
10	ATOM 2241	CZ	PHE	375	57.202	69.659	12.064	1.00	37.61
	ATOM 2242	Ċ.	PHE	375	52.982	67.917	9.664	1.00	28.21
	ATOM 2243	0	PHE	37 <i>5</i>	52.775	68.135	10.857	1.00	28.91
	ATOM 2244	N	SER	376	52.260	68.589	8.742	1.00	28.48
15	ATOM 2245	H	SER	376	52.414	68.468	7.783	1.00	0.00
15	ATOM 2246	CA	SER	376	51.281	69.562	9.208	1.00	28.11
	ATOM 2247	СВ	SER	376	50.715	70.398	8.047	1.00	24.30
	ATOM 2248	OG	SER	376	49.984	69.740	7.034	1.00	26.36
	ATOM 2249 ATOM 2250	HG	SER	376	50.480	68.972	6.709	1.00	0.00
20	ATOM 2251	c	SER	376	50.124	68.979	9.960	1.00	30,03
20	ATOM 2252	0	SER	376	49.395	69.683	10.647	1.00	27.69
	ATOM 2253	N H	PHE	377	49.927	67.671	9.8 38	1.00	33.33
	ATOM 2254	CA	PHE	377	50.511	67.127	9.265	1.00	0.00
	ATOM 2255	CB	PHE	377	48.882	66.990	10. <i>5</i> 70	1.00	30.24
25	ATOM 2256	CG	PHE	377	48.276	65.883	9.691	1.00	21.45
	ATOM 2257	CDI	PHE PHE	377	47.359	66.445	8.647	1.00	21.79
	ATOM 2258	CD2	PHE	377	47.815	66.654	7.363	1.00	23.05
	ATOM 2259	CEI	PHE	377	46.040	66.714	8.961	1.00	22.87
	ATOM 2260	CE2	PHE	377	46.933	67.097	6.390	1.00	23.70
30		CZ	PHE	377 377	45.173	67.155	7.981	1.00	21.35
	ATOM 2262	c	PHE	377	45.613	67.337	6.688	1.00	16.45
	ATOM 2263	ŏ	PHE	377	49.407	66.381	11.860	1.00	30.42
	ATOM 2264	N	GLU	378	48.713 50.634	65.615	12.528	1.00	29.72
	ATOM 2265	H	GLU	378	51.151	66,675 67,389	12.279	1.00	30.87
35	ATOM 2266	CA	GLU	378	51.151	65.910	11.850	1.00	0.00
	ATOM 2267	CB	GLU	378	52.649	66.182	13.374 13.534	1.00	35.50
	ATOM 2268	CG	GLU	378	53.146	64.886	14.168	1.00 1.00	33.01
	ATOM 2269	CD	GLU	378	54.632	64.790	14.100	1.00	38.65 42.24
	ATOM 2270	OEI	GLU	378	55.130	63.668	14.315	1.00	45.42
40	ATOM 2271	0E2	GLU	378	55.292	65.826	14.210	1.00	51.72
	ATOM 2272	С	GLU	378	50.502	66.092	14.719	1.00	36.27
	ATOM 2273	0	GLU	378	50.366	65.153	15.483	1.00	39.55
	ATOM 2274	N	GLN	379	50.053	67.255	15.116	1.00	39.19
	ATOM 2275	H	GLN	379	50.149	68.039	14.543	1.00	0.00
45	ATOM 2276	CA	GLN	379	49.268	67.367	16.328	1.00	48.22
	ATOM 2277	CB	GLN	379	49.380	68.788	16.896	1.00	56.51
	ATOM 2278	CG	GLN	379	50.771	69.437	16.941	1.00	63.32
	ATOM 2279	CD	GLN	379	51.711	68.603	17.767	1.00	69.92
F0	ATOM 2280	OEI	GLN	379	51.480	68.345	18.938	1.00	74.88
50	ATOM 2281	NE2	GLN	. 379	52.809	68.101	17.225	1.00	72.82
	ATOM 2282	HE21	GLN	379	52.996	68.281	16.281	1.00	0.00
	ATOM 2283	HE22	GLN	379	53.371	67.564	17.826	1.00	0.00
	ATOM 2284	C	GLN	379	47.797	67.066	16.009	1.00	50.39
55	ATOM 2285	0	GLN	379	47.241	67.7 23	15.130	1.00	51.95
55	ATOM 2286	N	PRO	380	47.0 9 8	66.157	16.602	1.00	51.98
	ATOM 2287	CD	PRO	380	47.634	65.095	17.439	1.00	50.16
	ATOM 2288	CA	PRO	380	45.670	65.984	16.367	1.00	58.10
	ATOM 2289	СВ	PRO	380	45.261	64.815	17.243	1.00	53.72
60	ATOM 2290	CG	PRO	380	46.548	64.024	17.310	1.00	49.03
00	ATOM 2291	C	PRO	380	44.890	67.242	16.664	00.1	67.86
	ATOM 2292	0	PRO	380	44.736	67.733	17.776	1.00	70.34
	ATOM 2293	N	ASP	381	44.366	67.794	15.586	1.00	77.19
	ATOM 2294	H	ASP	381	44.538	67.356	14.721	1.00	0.00
	ATOM 2295	CA	ASP	186	43.596	69.021	15.617	1.00	83.19

								69.494	14.153	1.00	90.96
	ATOM	2296	CB		NSP	381	43.541	70.910	13.938	1.00	08.27
	ATOM		CG		LSP	381	43.014	71.746	14.843	1.00	11.43
	ATOM		ODI		A.S.P	381	43.143	71.167	12.846	1.00	102.13
	ATOM		QD2		A.S.P	381	42.481 42.223	68.750	16.235	1.00	83.61
5	ATON		С		ASP	381	41.197	69.227	15.766	1.00	86.07
•	ATON		0		ASP	381	42.096	67.981	17.316	1.00	80.88
	ATON	4 2302	N		GLY	382 382	42.906	67.764	17.831	1.00	0.00
	ATON		Н		GLY	382	40.780	67.478	17.697	1.00	78.25 75.15
	ATO:		CA		GLY GLY	382	40.321	66.448	16.669	1.00	79.53
10	ATO:		Č		GLY	382	40.458	65.247	16.841	1.00	69 .56
	ATO:		0		ARG	383	39.759	66.873	15.545	1.00 1.00	0.00
	ATO		N H		ARG	383	39.519	67.824	15.480	1.00	64.31
	ATO:		CA		ARG	383	39.629	66.003	14.381 13.173	1.00	67.34
٦.	ATO	M 2310	CB		ARG	383	39.071	66.738	13.414	1.00	69.84
15	ATO	M 2311	CG		ARG	383	37.832	67.576	13.993	1.00	75.22
	ATO	M 2312			ARG	383	36.696	66.741	14.194	1.00	78.64
	ATC	M 2313			ARG	383	35.525	67.573	13.929	1.00	0.00
	ATC	M 2314	H		ARG	383	35.556	68.516 67.087	14.732	1.00	79.29
20	ATC	M 2315	C2		ARG	383	34,408	67.925	14.847	1.00	80.15
2,0	ATO				ARG	383	33.341	68.873	14.534	1.00	0.00
	ATO			H11	ARG	383	33.413 32.487	67.594	15,250	1.00	0.00
		OM 231		H12	ARG	383	34,322	65.793	15.173	1.00	76.12
	AT	OM 231	9 N	H2	ARG	383	34.322 35.108	65.180	15.101	1.00	0.00
25	AT	OM 232	0 H	H21	ARG	383	33.469	65.461	15.576	1.00	0.00
		OM 232	1 H	H22	ARG	383	41.041	65.557	14.005	1.00	59.27
	AT	OM 232			ARG	383 383	41.896	66.385	13.710	1.00	63.51
	AT	OM 232			ARG	384	41.392	64.279	13.985	1.00	51.06 0.00
		OM 237			ALA ALA	384	40.809	63.578	14.357	1.00	43.97
30		OM 23		i.	ALA	384	42,711	63.902	13.518	1.00	39.32
		OM 23		CA CB	ALA	384	43.392	63.101	14.618	1.00	41.10
		OM 23		C	ALA	384	42.614	63.088	12.237	1.00	42.25
		FOM 23		Ö	ALA	384	41.666	62.329	12.029 11.349	1.00	30.93
				N	GLN	385	43.604	63.245	11.422	1.00	0.00
3 5		TOM 23		Н	GLN	385	44.264	63.968	10.258	1.00	26.81
				CA	GLN	385	43.794	62.325 62.733	9.034	1.00	23.54
		-		СВ	GLN	385	42,964	64.098	8.481	1.00	27.75
			334	CG	GLN	385	43.318	64,422	7.399	1.00	29.52
4		TOM 2		CD	GLN	385	42.355 41.375	65.120	7.586	1.00	30.67
-3	<u> </u>	TOM 2	336	OE1	GLN	385	42.598	63.875	6.223	1.00	29.71
		TOM 2	337	NE2	GLN	385 385	43.370	63.272	6.110	1.00	0.00
		TOM 2	338	HE21	GLN	385	42.007		5.478	1.00	0.00
		TOM 2	339	HE22	GLN	385		1	9.937	1.00	26.43
4	5	ATOM 2		C	GLN	385			10.363	1.00	26.69 25.50
			2341	0	GLN	386			9.197	1.00	0.00
			2342	N	MET MET	386		60.770		1.00 1.00	25.53
			2343	H	MET	386		61.417			22.77
_			2344	CA	MET	. 386		60.441			22.60
=	50	MOTA	2343	CG	MET	386		60.20			24.47
		MOTA	2340 2347	SD	MET	386	6 50.36	9 61.71	0 9.227 0 10.768		22.26
		MOTA	2341	CE	MET	38	6 50.29	7 62.58			
		MOTA MOTA	2340	č	MET	38					
		MOTA	2350	ŏ	MET	38		44 =4			
:	55	ATOM	7351	N	PRO	38		40.0			
		MOTA	2352	CD	PRO	38					
		ATOM	2252	CA	PRO	38				•	
		MOTA	2355	СВ	PRO	38					
	6 0	ATOM	2355	CG	PRO		87 48.1				0 24.30
	60	ATOM	2356	c	PRO		87 48.8			·	0 23.73
		MOTA	2357	Ō	PRO	-	87 49.8				
		MOTA	2358	N	THR	_	88 48.6				ο.00
		ATOM	2359	H	THR	3	188 47.9	, Jy			
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,									

	ATOM 23	60 CA	THR	388	49.646	58.448	3.685	1.00	23.26
	ATOM 23	61 CB	THR	388	49.037	57.083	4.100	1.00	26.15
	ATOM 23		THR	388	48.399	57.238	5.359	1.00	34.81
_	ATOM 23		THR	388	49.021	57.669	5.956	1.00	0.00
5	ATOM 23		THR	388	50.105	56.009	4.257	1:00	24.27
	ATOM 23		THR	388	50.062	58.416	2.233	1.00	19.72
	ATOM 23		THR	388	49.262	58.508	1.316	1.00	20.10
	ATOM 23		THR	389	51.357	58.279	2.016	1.00	20.14
10	ATOM 23		THR	389	51.974	58.285	2.771	1.00	0.00
10	ATOM 23 ATOM 23		THR	389	51.933	58.099	0.709	1.00	22.14
	ATOM 23		THR THR	389	53.304	58.762	0.775	1.00	22.05
	ATOM 23		THR	389 389	53.043 52.584	60.143 60.204	0.521	00.1	32.86
	ATOM 23		THR	389	54.305	58.266	-0.321 -0.223	1.00 1.00	0.00
15	ATOM 23		THR	389	51.981	56.611	0.446	1.00	26.26 27.05
	ATOM 23		THR	389	52.421	55.845	1.307	1.00	23.21
	ATOM 23	76 N	GLU	390	51.532	56.183	-0.742	1.00	23.67
	ATOM 23	77 H	GLU	390	51.265	56.821	-1.440	1.00	0.00
	ATOM 23		GLU	390	51.461	54.770	-1.017	1.00	21.51
2,0	ATOM 23		GLU	390	49.993	54.334	-0.902	1.00	27.17
	ATOM 23		GLU	390	49.716	54.258	0.612	1.00	35.02
	ATOM 23		GLU	390	48.453	53.567	100.1	1.00	38.42
	ATOM 23		GLU	390	47.644	53.290	0.129	1.00	50.83
25	ATOM 23		GLU	390	48.274	53.314	2.192	1.00	47.46
25	ATOM 23		GLU	390	52.032	54.416	-2.351	1.00	22.05
	ATOM 23		GLU ARG	390	52.148	55.236	-3.244	1.00	25.04
	ATOM 23		ARG	391 391	52.406 52.314	53.145	-2.449	1.00	20.72
	ATOM 23		ARG	391	52.871	52.6 13 52.4 72	-1.634 -3.655	1.00 1.00	0.00 20.99
30	ATOM 23		ARG	391	51.677	52.087	-3.633 -4.548	1.00	24.04
	ATOM 23		ARG	391	51.351	50.594	-4.650	1.00	27.32
	ATOM 23	91 CD	ARG	391	51.041	50.062	-6.066	1.00	27.94
	ATOM 23		ARG	391	49.890	49.159	-6.094	1.00	36.16
	ATOM 23		ARG	391	49.248	49.211	-5.356	1.00	0.00
35	ATOM 23		ARG	391	49.642	48.255	-7.068	1.00	38.21
	ATOM 23		ARG	391	50.302	48.156	-8.24 6	1.00	39.70
	ATOM 23		ARG	391	51.042	48.794	-8.455	1.00	0.00
	ATOM 23 ATOM 23		ARG	391	50.046	47.453	-8.910	1.00	0.00
40	ATOM 23		ARG ARG	391 391	48.749	47,269	-6.849	1.00	47.46
- 0	ATOM 24		ARG	391	48,279 48.561	47,209 46,597	-5.968	1.00	0.00
	ATOM 24		ARG	391	53.842	53.303	-7,567 -4,477	1.00 1.00	0.00 24.89
	ATOM 24		ARG	391	53.590	53.655	-5.630	1.00	24.46
	ATOM 24		VAL	392	54.998	53.634	-3.892	1.00	25.48
45	ATOM 24	04 H	VAL	392	55.287	53.159	-3.085	1.00	0.00
	ATOM 24		VAL	392	55.880	54.629	-4,494	1.00	21.84
	ATOM 24		VAL	392	56.630	55.422	-3.404	1.00	19.22
	ATOM 24		VAL	392	57.408	56.628	-3.936	00.1	11.40
E-0	ATOM 24		VAL	392	55.579	55.918	-2.436	1.00	18.72
50	ATOM 24	••	VAL	392	56.865	53.948	-5.392	1.00	24.52
	ATOM 24		VAL	392	57.628	53.121	-4.915	1.00	20.00
	ATOM 24		THR THR	393 393	56.890	54.255	-6.691	1.00	23.61
	ATOM 24		THR	393	56.183 57.958	54.810	-7.081	1.00	0.00
55	ATOM 24		THR	393	57.450	53.731 52.681	-7.506 -8.535	1.00 1.00	20.89
	ATOM 24		THR	393	56.238	53.135	-9.148	1.00	21.30 22.68
	ATOM 24		THR	393	56.027	52.485	-9.822	1.00	0.00
	ATOM 24		THR	393	57.220	51.333	-7.855	1.00	17.80
	ATOM 24		THR	393	58.633	54.850	-8.235	1.00	22.52
60	ATOM 24		THR	393	59.201	54.661	-9.303	1.00	22.98
	ATOM 24		LEU	394	58.605	56.072	-7.719	1.00	19.99
	ATOM Z		LEU	394	58.080	56.246	-6.913	1.00	0.00
	ATOM 24		LEU	394	59.379	57.138	-8.338	1.00	20.68
	ATOM 24	23 CB	LEU	394	59.097	58.483	-7.647	1.00	18.77

						•		1.00	17.38
		CG	LEU	394	57.683	59.073	-7.778		16.87
	ATOM 2424		LEU	394	57.642	60.395	-7.031		17.9B
	ATOM 2425		LEU	394	57.320	59.347	-9,243	• • • •	21.53
	ATOM 2425		LEU	394	60.851	56.789	-8.181		21.60
_	ATOM 2427	_	LEU	394	61.273	56.518	-7.073		24.17
5	ATOM 2428	0	THR	395	61.659	56.769	-9.243	1.00	0.00
	ATOM 2429	N	THR	395	61.325	56.899	-10.160	1.00	22.29
	ATOM 2430	H	THR	395	63.090	56.503	-9.136	1.00	19.28
	ATOM 2431	. CA		395	63.474	55.581	-10.303	1.00	
	ATOM 2432	CB	THR	395	63.062	56.260	-11.471	1,00	22.57
10	ATOM 2433	OGI	THR	395	63.298	55.742	-12.244	1.00	0.00
	ATOM 2434	HGI	THR		62.748	54.238	-10.343	1.00	15.70
	ATOM 2435	CGZ	THR	395 395	63.957	57.778	-9.146	1.00	23.98
	ATOM 2436	С	THR		65.167	57.754	-8.943	1.00	25.97
	ATOM 2437	0	THR	395	63.362	58.947	-9.392	1.00	23.31
15	ATOM 2438	И	ARG	396	62.385	58.998	-9.428	1.00	0.00
	ATOM 2439	н	ARG	396	64.097	60.187	-9.519	1.00	26.02
	ATOM 2440	CA	ARG	396	64.189	60.695	-10.936	1.00	28.35
	ATOM 2441	CB	ARG	396	64.986	59.799	-11.825	1.00	32.64
	ATOM 2442	CG	ARG	396		60.158	-13.248	1.00	43.18
20	ATOM 2443	CD	ARG	396	64.666 65.266	59.145	-14.075	1.00	51.29
7 -	ATOM 2444	NE	ARG	396		58.271	-14.157	1.00	0.00
	ATOM 2445	HE	ARG	396	64.830	59.385	-14.715	1.00	57.25
	ATOM 2446	CZ	ARG	396	66.403	58.338	-15.382	1.00	62.75
	ATOM 2447	NHI	ARG	396	66.982	57.441	-15.369	1.00	0.00
25	ATOM 2448	HHII	ARG	396	66.539	58.466	-15.885	1.00	0.00
	ATOM 2449	HH12	ARG	396	67.836	60.635	-14.722	1.00	55.06
	ATOM 2450	NH2	ARG	396	66.961	61.394	-14.247	1.00	0.00
	ATOM 2451	HH21	ARG	396	66.516	60.786	-15.219	1.00	0.00
	ATOM 2452	HH22	ARG	396	67.815	61.210	-8.805	1.00	25.66
30		С.	ARG	396	63.286	60.990	-8.488	1.00	25.00
50	ATOM 2454	0	ARG	396	62.133	62.377	-8.546	1.00	27.52
	ATOM 2455	N	CYS	397	63.840		-8.830	1.00	0.00
	ATOM 2456	Н	CYS	397	64.766	62.545 63.404	-7.849	1.00	25.75
	ATOM 2457	CA	CYS	397	63.114		-7.306	1.00	33.69
35			CYS	397	64.086	64.447	-6.027	1.00	39.19
٥.	ATOM 2459		CYS	397	65.267	63.936	-8.834	1.00	24.55
	ATOM 2460		CYS	397	62.170	64.045	-9.977	1.00	23.49
	ATOM 2461		CYS	397	62.531	64.263	-8.464	1.00	21.44
	ATOM 2462		PHE	398	60.945	64.386	-7.560	1.00	0.00
			PHE	398	60.638	64.172	-9.380	1.00	20.44
4	ATOM 2464	CA	PHE	398	60.072	65.094	-9.380 -9.454	1.00	22.28
			PHE	398	58.677	64.374	-10.409	1.00	21.70
	ATOM 246: ATOM 246:		PHE	398	57.644	65.009	-11.494	1.00	26.69
			PHE	398	58.011	65.812		1.00	18.84
			PHE	398	56.306	64.770	-10.186	1.00	27.09
4			PHE	398	57.052	66.367	-12.329 -11.028	1.00	24.85
	••••		PHE	398	55.358	65.320		1.00	25.47
	ATOM 247		PHE	398	55.716	66.121	-12.095	1.00	22.33
	ATOM 247	-	PHE	398	59.960	66.512	-8.872	1.00	22.03
	ATOM 24		PHE	. 398		66.853	-7.926	1.00	24.80
5	50 ATOM 24	73 0	TYR	399		67.377	-9.537	1.00	0.00
	ATOM 24	74 N	TYR	399	44.484	67.062	-10.144		25.89
	ATOM 24	75 H		399			-9.395	1.00	25.90
	ATOM 24	76 CA	TYR	399			-9.624	1.00	
	ATOM 24	77 CB	TYR	399			-8.397	1.00	27.28
	55 ATOM 24	78 CG	TYR				-8.293	1.00	25.30
	ATOM 24	179 CD1		399		·		1.00	33.32
	ATOM 24	\$80 CE1		39	48				30.60
	ATOM 2	481 CD:		39				1.00	31.42
	ATOM 2	482 CE		39					31.10
	60 ATOM 2	483 CZ	TYR	39					35.89
	ATOM 2	484 OH		39					
	ATOM 2	485 HH			**	40.00			
	ATOM 2	486 C	TYR	39	9 59.53				
	ATOM 2	2487 O	TYR		99 59.66	58 69.10	, -11.03		
	AIOM 2	.,,,							

	ATOM	2488	N	LEU	400	58.468	69.931	-10.000	1.00	24.40
	ATOM		H	LEU	400	58.337	70.072	-9 .041	1.00	0.00
	ATOM		CA	LEU	400	57.598	70.629	-10.934	1.00	31.95
5	ATOM		CB	LEU	400	56.310	71.032	-10.186	1.00	27.67
5	ATOM		CG	LEU	400	55.445	69.821	-9.832	1.00	25.95
	ATOM ATOM		CD1 CD2	Leu Leu	400	54.671	70.119	-8.568	1.00	26.52
	ATOM		, C	LEU	400 400	54,550 58,422	69.476	-11.022	1.00	24.92
	ATOM		· õ	LEU	400	59.402	71.858 72.088	-11.293 -10.612	1.00 1.00	36.21 44.83
10	ATOM		N	PHE	401	58.287	72.708	-12.295	1.00	36.07
	ATOM		H	PHE	401	57.660	72.565	-13.041	1.00	0.00
	ATOM	2499	ĆA	PHE	401	59.210	73.839	-12.269	1.00	33.57
	ATOM		CB	PHE	401	60.245	73.706	-13.407	1.00	32.22
	MOTA		CG	PHE	401	61.399	72.857	-12.928	1.00	33.80
15	ATOM		CD1	PHE	401	61.474	71.519	-13.260	1.00	32.31
	ATOM ATOM		CD2	PHE	401	62.416	73.423	-12.173	1.00	39.98
	ATOM		CE1	PHE	401	62.582	70.764	-12.892	1.00	33.77
	ATOM		CZ	PHE PHE	401 401	63.512 63.611	72.653	-11.790	1.00	40.60
20	ATOM		C	PHE	401	58.367	71.326 75.031	-12.167 -12.475	1.00 1.00	36.46 32.19
•	ATOM		ō	PHE	401	58.334	75.584	-13.558	1.00	29.45
	ATOM		N	PRO	402	57.638	75.512	-11.527	1.00	29.88
	ATOM	2510	CD	PRO	402	57.751	75.159	-10.125	1.00	27.11
	ATOM		CA	PRO	402	56.710	76.609	-11.755	1.00	31.83
25	ATOM		CB	PRO	402	56.137	76.946	-10.380	1.00	27.59
	ATOM		CG	PRO	402	57.161	76.376	-9.428	1.00	24.71
	ATOM		C	PRO	402	57.461	77.757	-12.427	1.00	37.62
	ATOM		O N	PRO	402 .		78.123	-12.070	1.00	38.17
30	ATOM		H	GLY GLY	403 403	56.770	78.302	-13.431	1.00	37.83
20	ATOM		CA	GLY	403	55.816 57.339	78.096 79.300	-13.558 -14.327	. 1.00 1.00	0.00 40.55
	ATOM		C	GLY	403	58.024	78.691	-15.553	1.00	39.74
	ATOM	2520	0	GLY	403	58.363	79.436	-16.458	1.00	40.77
	ATOM		N	HIS	404	58.258	77.372	-15.646	1.00	40.90
35	MOTA		H	HIS	404	57.856	76.702	-15.043	1.00	0.00
	ATOM		CA	HIS	404	59.038	76.818	-16.735	1.00	41.51
	MOTA MOTA		CB	HIS	404	60.391	76.313	-16.275	1.00	43.86
	ATOM		CG CD2	ZIH ZIH	404 404	61.057	77.377	-15.470	1.00	47.48
40	ATOM		ND1	HIS	404	62.077 60.694	78.167 77.798	-15.913 -14.262	1.00	46.94 50.29
	ATOM		HDI	HIS	404	59.973	7 7.436	-13.710	1.00	0.00
	ATOM		CEI	HIS	404	61.438	78.828	-13.970	1.00	47.33
	ATOM	2530	NE2	HIS	404 .	62.266	79.045	-14.963	1.00	49.11
	ATOM		HEZ	HIS	404	62.923	79.771	-14.979	1.00	0.00
45	ATOM		C	HIS	404	58.318	75.633	-17.302	1.00	41.65
	ATOM		0	HIS	404	58.804	75.085	-18.288	1.00	44.03
	ATOM		OT	HIS	404	57.304	75.238	-16.718	1.00	41.54
	ATOM ATOM		OH2 H1	WAT WAT	256	57.131	53.937	-16.157	1.00	21.86
50	ATOM		H2	WAT	256 256	57.956 56.715	53.989 53.157	-16.611 -16.559	1.00 1.00	0.00 0.00
-	ATOM		OH2	WAT	257	59.288	45.222	-12.720	1.00	24.45
	ATOM		Hl	WAT	257	59.678	44.463	-12.289	1.00	0.00
	ATOM	2540	H2	WAT	257	59.326	45.020	-13.638	1.00	0.00
	MOTA		OH2	WAT	258	61.365	66.988	-12.454	1.00	18.38
55	ATOM		Hi	WAT	258	61.282	66.754	-11.566	1.00	0.00
		2543	H2	WAT	258	61.920	66.336	-12.878	1.00	0.00
	ATOM		OH2	WAT	259	54.401	52.311	-15.488	1.00	26.12
	MOTA		HI	TAW	259	53.455	52.320	-15.423	1.00	0.00
60	MOTA MOTA		H2	TAW	259	54.685	52.959	-14.831	1.00	0.00
50	ATOM		OH2 H1	TAW TAW	260 260	52.948 53.471	45.165	-10.748	1.00	22.53 0.00
	MOTA		H2	WAT	260	53.471 52.622	44.927 46.039	-9.991 -10.552	1.00	0.00
	ATOM		OH2	WAT	261	39.932	72.422	-0.681	1.00	41,66
	MOTA		HI	WAT	261	40,131	72.039	0.168	1.00	0.00
			• •							

								1.00	0.00
	ATOM 2552	H2 V	TAY	261	39.184	71.954	-1.011		27.37
	ATOM 2553		VAT	262	40.595	65.620	4.462 5.270	1.00	0.00
	ATOM 2554		TAY	262	40.213	65.976 65.614	3.842	1.00	0.00
	ATOM 2555		TAY	262	39.866	63.723	-5.839	1.00	23.97
5	ATOM 2556		TAY	263	59.703 59.734	63.118	-5.108	1.00	0.00
	ATOM 2557		TAY	263	59.734 59.203	63.239	-6.512	1.00	0.00
	ATOM 2558		WAT	263 264	57.975	70,486	-7.257	1.00	26.26
	ATOM 2559	~	WAT WAT	264	57.886	69.546	-7.232	1.00	0.00
	ATOM 2560		WAT	264	58.580	70.685	-6.537	1.00	0.00 51.01
10	ATOM 2561 ATOM 2562		WAT	265	49.889	74.051	7.407	1.00 1.00	0.00
	ATOM 2562 ATOM 2563		WAT	265	49.381	73.658	6.713 .	1.00	0.00
	ATOM 2564	H2	WAT	265	49.717	74.986	7.331 -12.629	1.00	54.87
	ATOM 2565	OH2	WAT	266	55.224	73.467	-12.029	1.00	0.00
15	ATOM 2566	HI	WAT	266	56.050	73.568 73.889	-13.488	1.00	0.00
	ATOM 2567	H2	WAT	266	55.324	72.666	-15,238	1.00	33.22
	ATOM 2568	OH2	WAT.	267	57.220 57.189	73.255	-16.021	1.00	0.00
	ATOM 2569	HI	WAT	267	56.964	71.837	-15.606	1.00	0.00
	ATOM 2570	H2	WAT	267 268	35.858	66.670	-2.607	1.00	29.58
20	ATOM 2571	OH2	WAT	268	36.152	66.629	-1.699	1,00	0.00
•	ATOM 2572	HI	WAT WAT	268	35.860	67.587	-2.844	1.00	0.00 49.48
	ATOM 2573	H2 OH2	TAW	269	48.789	71.281	-21.710	1.00	0.00
	ATOM 2574 ATOM 2575	HI	WAT	269	47.897	71.644	-21.837	1.00 1.00	0.00
אר	ATOM 2576	H2	WAT	269	49.355	71.928	-22.109	1.00	23.58
25	ATOM 2577	OH2	TAW	270	59.440	63.444	-17.067 -16.814	1.00	0.00
	ATOM 2578	H1	WAT	270	59.959	62.711	-10.014	1.00	0.00
	ATOM 2579	H2	WAT	270	58.513	63.193 44.941	-15.001	1.00	59.67
	ATOM 2580	OH2	TAW	271	48.923	44.016	-14.752	1.00	0.00
30	ATOM 2581	HI	WAT	271	48.905 49.386	44.932	-15.826	1.00	0.00
_	ATOM 2582	H2	TAW	271	44,435	59.093	9.639	1.00	24.08
	ATOM 2583	OHZ	TAW	272 272	44.989	59.185	8.877	1.00	0.00
	ATOM 2584	Hl	WAT	272	44.990	58.678	10.300	1.00	0.00
	ATOM 2585	H2 OH2	WAT WAT	273	53.920	52.043	-8.038	1.00	23,85 0.00
35	ATOM 2586 ATOM 2587	HI	WAT	273	54.603	52.710	-8.021	1.00 1.00	0.00
	ATOM 2588	H2	WAT	273	54.232	51.362	-8,608	1.00	38.92
	ATOM 2589		WAT	274	62.871	68.183	-1,698 -1,909	1.00	0.00
	ATOM 2590		WAT	274	62.632	67.292 68.466	-2.393	1.00	0.00
4(H2	TAW	274	63.467	70.044	-23.874	1.00	49.11
-	ATOM 2597	OH2	TAW	275	46.942 47.058	69.784	-24.775	1.00	0.00
	ATOM 259		WAT	275 275	47.406	69.414	-23.326	1.00	0.00
	ATOM 2594		TAW TAW	276	50.771	63.408	17.889	1.00	54.39
	ATOM 259		WAT	276	50.872	64.350	17.946	1.00	0.00 0.00
4			WAT	276	50.541	63.304	16.967	1.00	27.42
	ATOM 259 ATOM 259		TAW	277	45.555	65.749	12.972	1.00 1.00	0.00
	ATOM 259 ATOM 259	-	WAT	277	46.291	65.423	12.436 12.518	1.00	0.00
	ATOM 260		TAW	277	44.772	65.498	-11.771	1.00	27.22
5	O ATOM 260		TAW	278	57.066	46.788 46.154	-12.204	1.00	0.00
	ATOM 260		WAT	278	56.509		-12.005	1.00	0.00
	ATOM 260	3 H2	WAT	278	56.806		-29,411	1.00	47.42
	ATOM 26)4 QH2	TAW	279			-28.494	1.00	0.00
	ATOM 26	35 HI	WAT	279			-29.571	1.00	0.00
9	55 ATOM 26	06 H2	WAT	279 280			-27.639	1.00	73.43
	ATOM 26		WAT	280			-28.072	1.00	0.00
	ATOM 26	08 HI	TAW TAW	280			-27.145	1.00	0.00
	ATOM 26	09 H2		281		400	-26.241	1.00	45.60
	ATOM 26	10 OH2	WAT	281		5 51.280	-26.806	1.00	0.00
	60 ATOM 26	11 H2	WAT	281					51.46
	ATOM 26	12 612 13 OH2		283	2 40.35				0.00
	ATOM 2	513 OH1 514 H1	WAT	28	2 39.83				
	ATOM 2	615 H2	TAW	28	2 41.23	7 58.816	-25.588	1.00	7.40
	A10.4. 2								

·- 83 -

	ATOM	2616	OH2	WAT	283	59.582	57.507	-22.297		***
	ATOM	2617	HI	WAT	283	59.205	58.060		1.00	50.06
	ATOM	2618	H2	WAT	283	60.461		-22.973	1.00	0.00
	MOTA	2619	OH2	WAT	284		57.302	-22.574	1.00	0.00
5	ATOM		Hi	WAT		62.787	66.431	-21.929	1.00	6 4.67
	ATOM		H2		284	62.786	67.378	-21.776	1.00	0.00
	ATOM			WAT	284	63.310	66.298	-22.709	1.00	0.00
			OH2	WAT	285	42.178	68.676	-21.635	1.00	47.92
	ATOM		HI	WAT	285	42.226	68.152	-20.834	1.00	0.00
3.0	ATOM		H2	WAT	285	41_528	68.220	-22.154	1.00	0.00
10	MOTA		OH2	WAT	286	59.860	64.459	-20.626	1.00	45.90
	MOTA.		HI	WAT	286	59.529	63.613	-20.882	00.1	0.00
	MOTA	2627	HΣ	WAT	286	60.034	64.935	-21.429		
	ATOM	2628	OH2	WAT	287	38.592	60.429		1.00	0.00
	ATOM	2629	HI	WAT	287	37.765		-19.380	1.00	45.23
15	ATOM		H2	WAT	287		59.995	-19.240	1.00	0.00
	ATOM	2631	OH2	WAT		38.339	61.266	-19.751	1.00	0.00
	ATOM		Hi		288	49.737	64.079	-19.712	1.00	31.09
		2633		WAT	288	49.889	64.551	-20.540	1.00	0.00
			H2	WAT	288	48.791	63.991	-19.646	1.00	0.00
20	ATOM		OH2	WAT	289	45.077	74.284	-19.985	1.00	58.31
20	ATOM		HI	WAT	289	44.803	75.186	-19.856	1.00	0.00
	MOTA		H2	WAT	289	45.225	73.984	-19.086	1.00	0.00
		2637	OHZ	WAT	290	36.463	68.596	-18.372	1.00	41.38
		2638	H1	WAT	290	36.206	68.878	-19.244	1.00	
	ATOM	2639	H2	WAT	290	37.302	69.021	-18.240		0.00
25	MOTA	2640	OH2	WAT	291	42_509			1.00	0.00
	ATOM	2641	HI	WAT	291	42.018	73.175	-18.360	1.00	30.40
	MOTA		H2	WAT	291		73.868	-17.902	1.00	0.00
	ATOM		OH2	WAT		41.926	72.928	-19.071	1.00	0.00
	ATOM		Hi		292	52.772	54.057	-17.864	1.00	34.00
30	ATOM			WAT	292	52.675	54.050	-16.919	1.00	0.00
50			H2	WAT	292	53.709	54.139	-17.997	1.00	0.00
	MOTA		OH2	WAT	293	58.499	51.544	-17.264	1.00	39.19
	ATOM		HI	WAT	293	58.896	50.712	-17.042	1.00	0.00
	ATOM		H2	WAT	293	59.231	52.121	-17.413	1.00	0.00
3.5	MOTA		OH2	WAT	294	55.293	76.832	-15.365	1.00	77.96
35	ATOM		HI	WAT	294	55.586	76.689	-14.489	1.00	0.00
	ATOM	2651	H2	WAT	294	55.894	76.271	-15.896	1.00	0.00
	ATOM	2652	OH2	WAT	295	50.254	47.990	-11.950	1.00	
	ATOM	2653	HI	WAT	295	49.709	48.063			34.08
	ATOM	2654	H2	WAT	295	49.755		-12.721	1.00	0.00
40	ATOM		OH2	WAT	296		47.316	-11.486	1.00	0.00
		2656	HI	WAT	296	37.749	48.038	-8.897	1.00	50.16
	ATOM		H2	WAT		36.805	48.072	-8.734	1.00	0.00
	ATOM		OH2		296	37.815	47.302	-9.501	1.00	0.00
		2659		WAT	297	61.144	72.978	-8.832	1.00	30.80
45			HI	WAT	297	62.021	72.636	-8.821	1.00	0.00
30		2660	H2	WAT	297	61.120	73.515	-8.053	1.00	0.00
	MOTA		OH2	TAW	298	46.716	77.808	-7.102	1.00	36.58
		2662	HI	WAT	298	47.000	78.075	-6.217	1.00	0.00
		2663	H2	TAW	298	47.380	78.176	-7.649	1.00	0.00
	MOTA	2664	OH2	TAW	299	43.918	76.808	-6.081	1.00	35.53
50	ATOM	2665	Hl	WAT	. 299	43.850	77.750	-6.052	1.00	
	ATOM	2666	H2	WAT	299	44.809	76.650			0.00
	ATOM		OH2	WAT	300	60.882		-5.782	1.00	0.00
	ATOM		HI	WAT	300	60.547	61.010	-5.837	1.00	32.21
	ATOM		H2	WAT			60.543	-5.092	1.00	0.00
55		2670	OH2		300	60.943	61.933	-5.683	1.00	0.00
-				WAT	301	56.234	77.147	-5.325	1.00	45.88
		2671	HI	WAT	301	55.449	77.454	-4.859	1.00	9.00
		2672	H2	WAT	301	56.348	77.843	-5.97!	1.00	0.00
		2673	OH2	WAT	302	43.603	47.976	-4.116	1.00	45.63
		2674	HI	WAT	302	43. 96 9	48.651	-4.654	1.00	0.00
60	MOTA	2675	HZ	TAW	302	43.745	47.160	-4.601	1.00	0.00
		2676	OH2	WAT	303	41.712	55.660	-0.536	1.00	36.50
	ATOM		HI	WAT	303	41.333	54.851		1.00	
	MOTA		H2	WAT	303	42.325	55.359	-0.150		0.00
	ATOM		OH2	WAT	304			-1.193	1.00	0.00
			V.12	447	304	51.729	51.156	-0.590	1.00	72.02

	ATOM 2680	ні '	WAT	304	52.459	50.567	-0.423	1.00	0.00
	ATOM 2681		WAT	304	51.363	51.290	0.284	1.00	0.00 70.30
	ATOM 2682		TAW	305	44.576	76.180	0.284	1.00 1.00	0.00
	ATOM 2683	H1	TAW	305	44.696	75.258	0.070 -0.493	1.00	0.00
5	ATOM 2684		WAT	305	44.178	76.553 54.669	0.203	1.00	39.19
	ATOM 2685		WAT	306	38.913	55.452	-0.255	1.00	0.00
	ATOM 2686		WAT	306	39.203 38.207	54.284	-0.306	1.00	0.00
	ATOM 2687		TAW	306 307	42.134	58.338	1.150	1.00	29.79
	ATOM 2688	OH2 H1	WAT WAT	307	41.312	57.982	1.511	1.00	0.00
10	ATOM 2689 ATOM 2690	H2	WAT	307	42.564	57.551	0.838	1.00	0.00
	ATOM 2691	OH2	WAT	308	56.648	60.941	0.737	1.00	38.97 0.00
	ATOM 2692	Hl	WAT	308	55.700	60.977	0.666	1.00 1.00	0.00
	ATOM 2693	H2	WAT	308	56.928	61.839	0.583 9.192	1.00	48.96
15	ATOM 2694	OH2	WAT	309	45,030	48.554 47.651	9.474	1.00	0.00
	ATOM 2695	Hl	WAT	309	44.943 45.909	48.606	8.834	1.00	0.00
	ATOM 2696	H2	WAT	309 310	43.509	59.650	10.888	1.00	32.65
	ATOM 2697	OH2	WAT WAT	310	41.965	59.981	11.702	1.00	0.00
~ ^	ATOM 2698	H1 H2	WAT	310	41.171	60.430	10.534	1.00	0.00
20	ATOM 2699 ATOM 2700	OH2	WAT	311	30.678	62.812	10.599	1.00	48.30
	ATOM 2701	HI	WAT	311	31.519	63.059	10.280	1.00	0.00 0.00
	ATOM 2702	H2	WAT	311	30.787	61.904	10.876	1,00	49.62
	ATOM 2703	OH2	WAT	312	44.035	51.425	12.296 11.383	1.00	0.00
25	ATOM 2704	Hl	WAT	312	43.759	51.313	12.653	1.00	0.00
	ATOM 2705	H2	WAT	312	43.889	50.557 69.483	13.408	1.00	45.11
	ATOM 2706	OH2	WAT	313 313	53.084 53.666	68.732	13.409	1.00	0.00
	ATOM 2707	HI	TAW TAW	313	52.885	69.526	12.474	1.00	0.00
20	ATOM 2708 ATOM 2709	H2 OH2	WAT	314	33.280	54.578	14.147	1.00	71.20
30	ATOM 2709 ATOM 2710	H!	WAT	314	32.487	54.073	14.271	1.00	0.00
	ATOM 2711	H2	TAW	314	33.372	54.689	13.208	1.00 1.00	30.28
	ATOM 2712	OH2	WAT	315	60.509	60.787	-18.332	1.00	0.00
	ATOM 2713	Hi	TAW	315	60.849	61.538	-18.810 -17.565	1.00	0.00
35	ATOM 2714	H2	TAW	315	60.079	61.112 51.291	10.254	1.00	49.82
	ATOM 2715	OH2	WAT	316	36.436 36.114	50.786	9.515	1.00	0.00
	ATOM 2716	HI	WAT WAT	316 316	35.650	51.562	10.727	1.00	0.00
	ATOM 2717	H2 OH2	WAT	317	47.543	66.402	-29.189	1.00	49.95
4.0	ATOM 2718 ATOM 2719	HI	WAT	317	46,808	66.985	-29.300	1.00	0.00 0.00
40	ATOM 2719	H2	WAT	317	48.149	66.611	-29.900	1.00 1.00	55.32
	ATOM 2721	OH2	WAT	318	39.908	61.653	-21.569 -20.946	1.00	0.00
	ATOM 2722	Hi	TAW	318	39.811	62.356	-20.940 -21.097	1.00	0.00
	ATOM 2723	H2	TAW	318	40.435 43.648	61.009 51.317	-20.053	1.00	55.22
45	ATOM 2724	OH2	TAW	319 319	44.217	51.632	-19.345	1.00	0.00
	ATOM 2725	HI	TAW TAW	319	42.798	51.245	-19.643	1.00	0.00
	ATOM 2726		WAT	320	42.904	66.185	-19.404	1.00	43.44
	ATOM 2727 ATOM 2728		TAW	320	43.844	66.182	-19.470	1.00	0.00 0.00
5			WAT	. 320	42.797	66.244	-18.456	1.00 1.00	47.88
٠,	ATOM 273		TAW	321	52,576	73.792	-19.312	1.00	0.00
	ATOM 273		WAT	321	52.248	73.438	-18.497 -19.932	1.00	0.00
	ATOM 273	2 H2	WAT	321	51.924	73.486 50.185	-16.806	1.00	60.71
	ATOM 273	3 OH2	WAT	322	61.556	49.697	-16.824	1.00	0.00
5	5 ATOM 273		WAT	322	60.747 62.307	49.596	-16.932	1.00	0.00
	ATOM 273	5 H2	TAW	322 323	24.851	56.075	-3.153	1.00	38.05
	ATOM 273	6 OH2	WAT WAT	323 323	25.114		-2.507	1.00	0.00
	ATOM 273	7 H1 18 H2	WAT	323			-3,916	1.00	0.00
_	ATOM 273			324	30.036	71.409	-2.744	1.00	45.31 0.00
6	0 ATOM 273 ATOM 274		WAT	324	29.383		-3.022	1.00	0.00
	ATOM 27		WAT	324	29.785		-1.852 0.558	1.00	57.23
	ATOM 27	42 OH2		325			-0.294	1.00	0.00
	ATOM 27	43 H1	WAT	325	33.528	17,404			

	ATOM	2744	H2	WAT	325	32.600	60 047	0.700		
	ATOM	2745	OH2	WAT	326		80.047	0.708	1.00	0.00
	ATOM		H!	WAT		35.907	47.459	2.721	1.00	52.14
	ATOM				326	35.224	48.120	2.665	1.00	0.00
5	ATOM		H2	WAT	326	36.753	47.864	2.620	1.00	0.00
		2748	OH2	WAT	327	54.215	72.016	6.546	1.00	52.02
	ATOM	2749	H1	WAT	327	55.027	71.530	5.405	1.00	0.00
	ATOM	2750	H2	WAT	327	54.516	72.859	6.883	1.00	0.00
	ATOM	2751	OH2	WAT	328	41.269	52.487	1.122	1.00	51.87
4.0	ATOM	2752	H1	WAT	328	40.694	51.781	1.440	1.00	0.00
10	ATOM	2753	H2	WAT	328	42.127	52.127	1.259	1.00	0.00
	ATOM	2754	OH2	WAT	329	34.066	58.806	13.164	1.00	55.71
	ATOM	2755	Hi	WAT	329	34.724	59.474	13.292	1.00	0.00
	ATOM	2756	H2	WAT	329	34.564	58.010	12.945	1.00	0.00
	ATOM	2757	OH2	WAT	330	41.816	52.756	13.918	1.00	44.03
15	ATOM	2758	Hi	WAT	330	42.281	53.408	13.395	1.00	0.00
	ATOM	2759	H2	WAT	330	42,525	52.335	14.395	1.00	0.00
	ATOM	2760	OH2	WAT	331	39.370	62.098	14.302	1.00	54.70
	MOTA	2761	HI	TAW	331	38.736	62.661	14.727	1.00	
	ATOM	2762	H2	WAT	331	39.761	62.569	13.574		0.00
20	ATOM	2763	OH2	WAT	332	50.309	69.365		1.00	0.00
•	ATOM	2764	HI	WAT	332			13.364	1.00	54.23
	ATOM	2765	H2			50.055	69.719	12.508	1.00	0.00
	ATOM	2766		WAT	332	51.043	69.910	13.645	1.00	0.00
	ATOM		OH2	TAW	333	40.562	55.451	15.773	1.00	61.39
25	ATOM	2767	Hi	WAT	333	39.723	55.080	16.041	1.00	0.00
23	VIOW	2768	H2	TAW	333	40.748	55.017	14.937	1.00	0.00

The following abbreviations are used in Table B.

"Atom type" refers to the element whose
coordinates are measured. The first letter in the
column defines the element.

30 "X, Y, Z" crystallographically define the atomic position of the element measured.

"B" is a thermal factor that measures movement of the atom around its atomic center.

Atoms numbered 153-158 (Lys-146) and 184-189 (Ser-149) were modeled as Ala residues.

Atoms numbered 1487-1534 and designated "Ald" in the column titled "Residue" are Cys-285 bound to the tetrapeptide aldehyde inhibitor.

Structure coordinates for ICE according to Table B
40 may be modified from this original set by mathematical
manipulation. Such manipulations include, but are not
limited to, crystallographic permutations of the raw
structure coordinates, fractionalization of the raw
structure coordinates, integer additions or

45 subtractions to sets of the raw structure coordinates,

inversion of the raw structure coordinates, and any combination of the above.

- 87 -

SEQUENCE LISTING

	(1) GENERAL INFORMATION:	
5	/// Appropries	
	(i) APPLICANT:	
	(A) NAME: Vertex Pharmaceuticals, Inc.	
	(B) STREET: 40 Allston Street	
10	(C) CITY: Cambridge	
	(D) STATE: Massachusetts	
	(E) COUNTRY: United States of America (F) POSTAL CODE (ZIP): 02139	
	(G) TELEPHONE: 617-576-3111	
	(H) TELEFAX: 617-576-2109	
15	, 1001AX: 01/-3/6-2109	
	(ii) TITLE OF INVENTION: CRYSTAL STRUCTURE AND MUTANTS OF	
	INTERLEUKIN-1 BETA CONVERTING ENZYME	
	(iii) NUMBER OF SEQUENCES: 1	
20	11. 1	
	(iv) COMPUTER READABLE FORM:	
	(A) MEDIUM TYPE: Floppy disk	
	(B) COMPUTER: IBM PC compatible	
25	(C) OPERATING SYSTEM: PC-DOS/MS-DOS	
	(D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)	
	(vi) PRIOR APPLICATION DATA:	
	(A) APPLICATION NUMBER: US 08/261,582	
	(B) FILING DATE: 17-JUN-1994	
30	The state of the s	
	(2) INFORMATION FOR SEQ ID NO: 1:	
	(i) SEQUENCE CHARACTERISTICS:	
35	(A) LENGTH: 404 amino acids	
	(B) TYPE: amino acid	
	(C) STRANDEDNESS:	
	(D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: protein	
	(iii) HYPOTHETICAL: NO	
45		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
	Mak 23a Nam 1sta 25-1 a	
	Met Ala Asp Lys Val Leu Lys Glu Lys Arg Lys Leu Phe Ile Arg Ser	
50	1 5 10 15	
	Met Gly Gly Gly The Tle New City	
	Met Gly Glu Gly Thr Ile Asn Gly Leu Leu Asp Glu Leu Leu Gln Thr	
	20 25 30	

- 88 -

	Arg Val Leu Asn Lys Glu Glu Met Glu Lys Val Lys Arg Glu Asn Ala 35 40 45
5	Thr Val Met Asp Lys Thr Arg Ala Leu Ile Asp Ser Val Ile Pro Lys 50 55 60
	Gly Ala Gln Ala Cys Gln Ile Cys Ile Thr Tyr Ile Cys Glu Glu Asp 65 70 75 80
10	Ser Tyr Leu Ala Gly Thr Leu Gly Leu Ser Ala Asp Gln Thr Ser Gly 85 90 95
	Asn Tyr Leu Asn Met Gln Asp Ser Gln Gly Val Leu Ser Ser Phe Pro 100 105 110
	Ala Pro Gln Ala Val Gln Asp Asn Pro Ala Met Pro Thr Ser Ser Gly 115 120 125
20	Ser Glu Gly Asn Val Lys Leu Cys Ser Leu Glu Glu Ala Gln Arg Ile 130 135 140
	Trp Lys Gln Lys Ser Ala Glu Ile Tyr Pro Ile Met Asp Lys Ser Ser 145 150 155 160
25	Arg Thr Arg Leu Ala Leu Ile Ile Cys Asn Glu Glu Phe Asp Ser Ile 165 170 175
	Pro Arg Arg Thr Gly Ala Glu Val Asp Ile Thr Gly Met Thr Met Leu 180 185 190
30	Leu Gln Asn Leu Gly Tyr Ser Val Asp Val Lys Lys Asn Leu Thr Ala 195 200 205
35	Ser Asp Met Thr Thr Glu Leu Glu Ala Phe Ala His Arg Pro Glu His 210 215 220
	Lys Thr Ser Asp Ser Thr Phe Leu Val Phe Met Ser His Gly Ile Arg 225 230 235 240
40	Glu Gly Ile Cys Gly Lys Lys His Ser Glu Gln Val Pro Asp Ile Leu 255
	Gln Leu Asn Ala Ile Phe Asn Met Leu Asn Thr Lys Asn Cys Pro Ser 260 265 270
45	Leu Lys Asp Lys Pro Lys Val Ile Ile Ile Gln Ala Cys Arg Gly Asp 275 280 285
50	Ser Pro Gly Val Val Trp Phe Lys Asp Ser Val Gly Val Ser Gly Ass 290 295 300
	Leu Ser Leu Pro Thr Thr Glu Glu Phe Glu Asp Asp Ala Ile Lys Ly:

WO 95/35367 PCT/US95/07619

- 89 -

	Ala	His	Ile	Glu	Lys 325	Asp	Phe	Ile	Ala	Phe 330	Cys	Ser	Ser	Thr	Pro 335	Asp
5	Asn	Val	Ser	Trp 340	Arg	His	Pro	Thr	Met 345	Gly	Ser	Val	Phe	Ile 350	Gly	Arg
10	Leu	Ile	Glu 355	His	Met	Gln	Glu	Tyr 360	Ala	Cys	Ser	Cys	Asp 365	Val	Glu	Glu
	Ile	Phe 370	Arg	Lys	Val	Arg	Phe 375	Ser	Phe	Glu	Gln	Pro 380	Asp	Gly	Arg	Ala
15	Gln 385	Met	Pro	Thr	Thr	Glu 390	Arg	Val	Thr	Leu	Thr 395	Arg	Cys	Phe	Tyr	Leu 400
	Phe	Pro	Gly	His												

PCT/US95/07619 WO 95/35367

- 90 -

CLAIMS

We claim:

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- 1. An interleukin-16 converting enzyme crystal, wherein said crystal has tetragonal space group symmetry $P4_32_12$.
- 2. The interleukin-1ß converting enzyme crystal according to claim 1, wherein said crystal comprises rectangular shaped unit cells, each unit cell having the dimensions $a=b=65\pm5\text{\AA}$, and $c=162\pm5\text{\AA}$.
- 3. The interleukin-1ß converting enzyme crystal according to claim 1, wherein said enzyme is a tetramer.
 - 4. The interleukin-18 converting enzyme crystal according to claim 3, wherein said tetramer comprises two adjacent pl0 subunits contacted by two p20 subunits, said p10 subunits interacting across the two-fold axis of said crystal.
 - 5. The interleukin-18 converting enzyme crystal according to claim 4, wherein said enzyme is characterized by an active site moiety characterized by at least amino acids 173, 176, 177, 178, 179, 180, 236, 237, 238, 239, 244, 248, 283, 284, 285, 290, 338, 339, 340, 341, 342, 343, 344, 345, 348, 352, 381 and 383 of SEQ. ID NO:1.
- 6. The interleukin-1ß converting enzyme crystal according to claim 5, wherein said active site moiety comprises amino acids from said pl0 and p20 subunits.

WO 95/35367 PCT/US95/07619

- 91 -

- 7. The interleukin-1ß converting enzyme crystal according to claim 4, wherein said enzyme is characterized by an accessory binding site moiety characterized by at least amino acids 150, 151, 240, 259, 267, 268, 274, 291, 292, 293, 294, 295, 296, 297, 317, 318, 319, 320, 321, 322, 323, 324, 325, 327, 334, 335, 367, 371, 374, 375, 377, 378, 380, 382, 384, 386, 388, 389, 390, 391, 392, 393, 394, 395 and 396 of SEQ. ID NO:1.
- 8. The interleukin-1ß converting enzyme crystal according to claim 7, wherein said accessory binding site moiety comprises amino acids adjacent to said two-fold axis according to Table A.
- 9. The interleukin-1ß converting enzyme crystal according to any one of claims 1 to 8, wherein said enzyme is characterized by structure coordinates according to Table B.

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- 10. A heavy atom derivative of a crystal, said crystal being selected from the group consisting of crystals of ICE, crystals of ICE mutants, crystals of ICE homologues or crystals of co-complexes of ICE.
- 11. The heavy atom derivative according to claim 10, wherein said derivative is formed by the reaction of said crystal with a compound selected from the group consisting of thimerosal, gold thiomalate, uranyl acetate and lead chloride.
- 12. The use of the structure coordinates of interleukin-1ß converting enzyme, or portions thereof, to solve a crystal form of a mutant, homologue or co-complex of interleukin-1ß converting enzyme by molecular replacement.

WO 95/35367

- 92 -

13. The use of the structure coordinates of interleukin-1ß converting enzyme to computationally evaluate a chemical entity for associating with the active site or the accessory binding site of interleukin-1ß converting enzyme.

14. The use of the structure coordinates of interleukin-1ß converting enzyme to design a compound capable of associating with the active site or the accessory binding site of interleukin-1ß converting enzyme.

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- 15. The use of the structure coordinates of interleukin-1ß converting enzyme according to claim 13 or 14, wherein a compound that is characterized by the chemical entity that associates with said active site or said accessory binding site, is an inhibitor of interleukin-1ß converting enzyme.
- 16. The use of the structure coordinates according to claim 15, wherein said inhibitor is a non-competitive or uncompetitive inhibitor of interleukin-16 converting enzyme.
- 17. The use of the structure coordinates of interleukin-1ß converting enzyme to determine the orientation of ligands in the active site or in the accessory binding site of interleukin-1ß converting enzyme.
- 18. The use of the structure coordinates of interleukin-18 converting enzyme to identify an intermediate in a chemical reaction between said enzyme and a compound which is an ICE substrate or an ICE inhibitor.

WO 95/35367 PCT/US95/07619

- 93 -

19. The use of the structure coordinates of interleukin-1ß converting enzyme according to any one of claims 12 to 14 or 17 to 18, wherein said structure coordinates are according to Table B.

The use of the structure coordinates of interleukin-1ß converting enzyme according to claim 15, wherein said structure coordinates are according to Table B.

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21. The use of the structure coordinates of interleukin-1ß converting enzyme according to claim 16, wherein said structure coordinates are according to Table B.

- 22. An interleukin-16 converting enzyme, wherein one or more of the amino acids in the active site or in the accessory binding site are replaced by one or more amino acids selected from the group consisting of naturally occurring amino acids, unnatural amino acids, selenocysteine and selenomethionine.
- 23. The interleukin-1ß converting enzyme according to claim 22, wherein a hydrophilic or hydrophobic amino acid residue in said active site or said accessory binding site is replaced.
- 24. The interleukin-1ß converting enzyme according to claim 22, wherein said active site amino acid is selected from the group consisting of amino acids 173, 176, 177, 178, 179, 180, 236, 237, 238, 239, 244, 248, 283, 284, 285, 290, 338, 339, 340, 341, 342, 343, 345, 348, 352, 381 and 383 of SEQ. ID NO:1.

WO 95/35367 PCT/US95/07619

· - 94 -

- 25. The interleukin-1ß converting enzyme according to claim 22, wherein said accessory binding site amino acid is selected from the group consisting of amino acids 150, 151, 240, 259, 267, 268, 274, 291, 292, 293, 294, 295, 296, 297, 317, 318, 319, 320, 321, 322, 323, 324, 325, 327, 334, 335, 367, 371, 374, 375, 377, 378, 380, 382, 384, 386, 388, 389, 390, 391, 392, 393, 394, 395 and 396 of SEQ. ID NO:1.
- 26. The interleukin-18 converting enzyme

 according to claim 22, wherein at least one cysteine
 amino acid is replaced by an amino acid selected from
 the group consisting of selenocysteine or
 selenomethionine.

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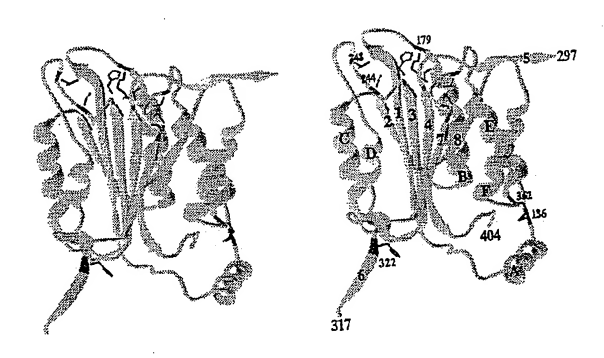
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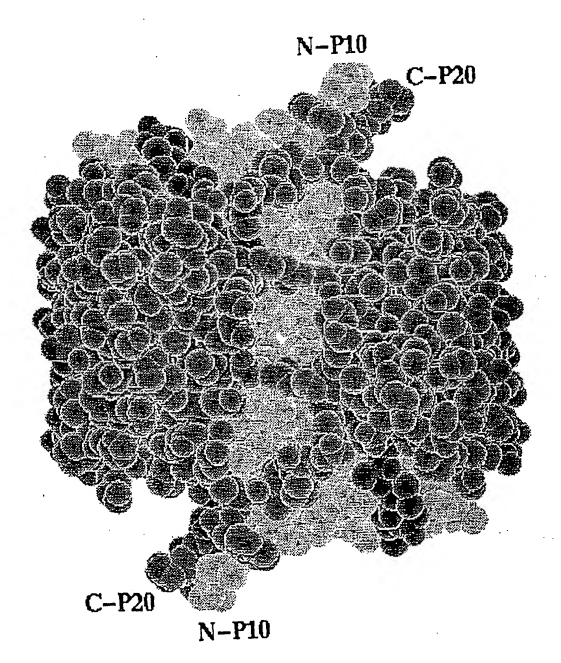
- 27. The interleukin-1ß converting enzyme according to claim 22, wherein at least one methionine amino acid is replaced by an amino acid selected from the group consisting of selenocysteine or selenomethionine.
 - 28. The interleukin-1ß converting enzyme according to any one of claims 22 to 27, wherein said enzyme is in crystalline form.
 - 29. The interleukin-16 converting enzyme according to claim 22, wherein said enzyme is characterized by increased stability to subunit dissociation.
 - 30. The interleukin-1ß converting enzyme according to claim 22, said enzyme having higher specific activity than the wild-type enzyme.

- 31. The interleukin-1ß converting enzyme according to claim 22, said enzyme having altered substrate specificity.
- 32. The use of an interleukin-1ß converting enzyme according to claim 22 to determine binding interactions between a chemical compound and the enzyme.
- 33. An interleukin-1ß converting enzyme,
 wherein at least one amino acid residue on, at or near
 the surface of said enzyme is replaced, resulting in an
 altered surface charge of one or more charge units.

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2/3



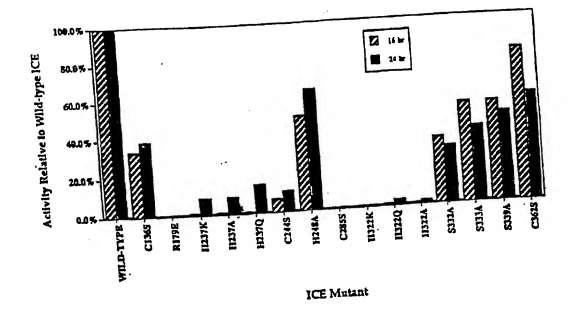


FIGURE 3

INTERNATIONAL SEARCH REPORT

Intern val Application No PCT/US 95/07619

	•		C1/02 32/0/913
L CLASSIF	C12N9/64 C12Q1/37		
			•
of gainage	International Patrat Classification (IPC) or to both national cl	assification and IPC	
	SEARCHED		
Ainimum do IPC 6	cumentation searched (dassification system followed by dassification fol	fication symbols)	
ocumentati:	on searched other than minimum documentation to the extent t	hat such documents are include	d in the fields searched
dectronic da	its base committed during the international search (name of data	base and, where practical, sear	rch terms used)
. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category '	Citation of document, with indication, where appropriate, of t	he relevant passages	Relevant to claim No.
X	JOURNAL OF CELLULAR BIOCHEMIST	RY,	1-9
	vol. 18D, 5 - 12 March 1994 page 148		
	J. THOMSON ET AL 'In vito following autoprocessing of active interbeta converting enzyme from an	leukin-1	
Y	recombinant precursor ¹ see abstract S 229		10-33
x	WO,A,94 00154 (MERCK & CO INC ANDREW D (US); MOLINEAUX SUSAN	;HOWARD M (US);	1-9
Y	TOCC) 6 January 1994 see the whole document		10-33
	Pr- vol dill	-/	
X Furt	ther documents are listed in the continuation of box C.	X Patent family me	unhers are listed in annex.
'A' docum	tegories of cited documents : ent defining the general state of the art which is not error to be of particular relevance	or priority date and a	shed after the international filing date not in conflict with the application but the principle or theory underlying the
E' earlier	document but published on or after the international	carmot be considered involve an inventive	lar relevance; the claimed invention d novel or cannot be considered to step when the document is taken alone
which citato "O" docum	is cited to establish the publication date of another m or other special reason (as specified) tent referring to an oral disclosure, use, exhibition or means	Y document of particul carnot be considere document is combin ments, such combin	lar relevance; the daimed invention d to involve an inventive step when the sed with one or more other such docu- ation being obvious to a person skilled
	ent published prior to the international filing date but than the priority date claimed		f the same patent family
	actual completion of the international search	Date of mailing of the 29. 11.	ne international search report
2	23 October 1995		
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	Authorized officer	
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Facc (+31-70) 340-3016	Van der	Schaal, C

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